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SUBSTITUTED ALKYLAMINE DERIVATIVES AND METHODS OF USE

This application claims the benefit of U.S. Provisional Application Nos. 60/261,339, filed January 12, 2001, and 60/323,764 filed September 19, 2001 which are hereby incorporated by reference.

FIELD OF THE INVENTION

This invention is in the field of pharmaceutical agents and specifically relates to compounds, compositions, uses and methods for treating cancer and angiogenesisrelated disorders.

BACKGROUND OF THE INVENTION

Protein kinases represent a large family of proteins which play a central role in the regulation of a wide variety of cellular processes, maintaining control over cellular function. A partial list of such kinases includes abl, Atk, bcr-abl, Blk, Brk, Btk, c-kit, c-met, c-src, CDK1, CDK2, CDK3, CDK4, CDK5, CDK6, CDK7, CDK8, CDK9, CDK10, cRaf1, CSF1R, CSK, EGFR, ErbB2, ErbB3, ErbB4, Erk, Fak, fes, FGFR1, FGFR2, FGFR3, FGFR4, FGFR5, Fdr. flt-1.

25 Fps, Frk, Fyn, Hck, IGF-1R, INS-R, Jak, KDR, Lck, Lyn, MEK, p38, PDGFR, PIK, PKC, PYK2, ros, tie, tie2, TRK, Yes, and Zap70. Inhibition of such kinases has become an important therapeutic target.

Certain diseases are known to be associated with

deregulated angiogenesis, for example ocular
neovascularization, such as retinopathies (including
diabetic retinopathy), age-related macular degeneration,
psoriasis, hemangioblastoma, hemangioma, arteriosclerosis,
inflammatory disease, such as a rheumatoid or rheumatic

inflammatory disease, especially arthritis (including
rheumatoid arthritis), or other chronic inflammatory
disorders, such as chronic asthma, arterial or post-

transplantational atherosclerosis, endometriosis, and neoplastic diseases, for example so-called solid tumors and liquid tumors (such as leukemias).

At the center of the network regulating the growth and 5 differentiation of the vascular system and its components, both during embryonic development and normal growth, and in a wide number of pathological anomalies and diseases, lies the angiogenic factor known as Vascular Endothelial Growth Factor"(VEGF; originally termed 'Vascular Permeability 10 Factor", VPF), along with its cellular receptors (see G.

Breier et al., Trends in Cell Biology, 6, 454-6 (1996)).

VEGF is a dimeric, disulfide-linked 46-kDa
glycoprotein related to "Platelet-Derived Growth Factor"
(PDGF); it is produced by normal cell lines and tumor cell

15 lines; is an endothelial cell-specific mitogen; shows angiogenic activity in in vivo test systems (e.g. rabbit cornea); is chemotactic for endothelial cells and monocytes; and induces plasminogen activators in endothelial cells, which are involved in the proteolytic degradation of

20 extracellular matrix during the formation of capillaries. A number of isoforms of VEGF are known, which show comparable biological activity, but differ in the type of cells that secrete them and in their heparin-binding capacity. In addition, there are other members of the VEGF family, such 25 as "Placenta Growth Factor" (PIGF) and VEGF-C.

VEGFR receptors (VEGFR) are transmembranous receptor tyrosine kinases. They are characterized by an extracellular domain with seven immunoglobulin-like domains and an intracellular tyrosine kinase domain. Various types of VEGF receptor are known, e.g. VEGFR-1 (also known as flt-1), VEGFR-2 (also known as KDR), and VEGFR-3.

A large number of human tumors, especially gliomas and carcinomas, express high levels of VEGF and its receptors. This has led to the hypothesis that the VEGF released by

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tumor cells stimulates the growth of blood capillaries and the proliferation of tumor endothelium in a paracrine manner and through the improved blood supply, accelerate tumor growth. Increased VEGF expression could explain the occurrence of cerebral edema in patients with glioma. Direct evidence of the role of VEGF as a tumor angiogenesis factor in vivo is shown in studies in which VEGF expression or VEGF activity was inhibited. This was achieved with anti-VEGF antibodies, with dominant-negative VEGFR-2 mutants which inhibited signal transduction, and with antisense-VEGF RNA techniques. All approaches led to a reduction in the growth of glioma cell lines or other tumor cell lines in vivo as a result of inhibited tumor angiogenesis.

Angiogenesis is regarded as an absolute prerequisite for tumors which grow beyond a diameter of about 1-2 mm; up to this limit, oxygen and nutrients may be supplied to the tumor cells by diffusion. Every tumor, regardless of its origin and its cause, is thus dependent on angiogenesis for its growth after it has reached a certain size.

Three principal mechanisms play an important part in the activity of angiogenesis inhibitors against tumors: 1) Inhibition of the growth of vessels, especially capillaries, into avascular resting tumors, with the result that there is no net tumor growth owing to the balance that is achieved between cell death and proliferation; 2) Prevention of the migration of tumor cells owing to the absence of blood flow to and from tumors; and 3) Inhibition of endothelial cell proliferation, thus avoiding the paracrine growth-stimulating effect exerted on the surrounding tissue by the endothelial cells which normally line the vessels. See R. Connell and J. Beebe, Exp. Opin. Ther. Patents, 11, 77-114 (2001).

VEGF's are unique in that they are the only angiogenic growth factors known to contribute to vascular

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hyperpermeability and the formation of edema. Indeed, vascular hyperpermeability and edema that is associated with the expression or administration of many other growth factors appears to be mediated via VEGF production.

Inflammatory cytokines stimulate VEGF production.

Hypoxia results in a marked upregulation of VEGF in numerous tissues, hence situations involving infarct, occlusion, ischemia, anemia, or circulatory impairment typically invoke VEGF/VPF-mediated responses. Vascular hyperpermeability, associated edema, altered transendothelial exchange and

associated edema, altered transendothelial exchange and macromolecular extravasation, which is often accompanied by diapedesis, can result in excessive matrix deposition, aberrant stromal proliferation, fibrosis, etc. Hence, VEGF-mediated hyperpermeability can significantly contribute to disorders with these etiologic features. As such, regulators of angiogenesis have become an important therapeutic target.

Schipper US patent 3,226,394, issued Dec. 28, 1965, describes anthranilamides as CNS depressants. Japanese patent JP2000256358 describes pyrazole derivatives that block the calcium release-activated calcium channel. EP application 9475000, published 6 October 1999, describes compounds as PGE2 antagonists. PCT publication WO96/41795, published 27 December 1996, describes benzamides as vasopressin antagonists. WO01/29009 describes aminopyridines as KDR inhibitors. WO01/30745 describes anthranilic acids as CGMP phosphodiesterase inhibitors. WO00/02851, published 20 Jan 2000 describes arvIsulfonylamnoaryl amides as quanylate cyclase activators.

W098/45268 describes nicotinamide derivatives as PDE4
30 inhibitors. W098/24771 describes benzamides as vasopressin antagonists.

US Patent No. 5,532,358, issued July 2, 1996, describes the preparation of 2-(cyclopropylamino)-N-(2-methoxy-4-methyl-3-pyridinyl)-3-pyridinecarboxamide as an

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intermediate for HIV inhibitors. Triazine-substituted amines are described for their aggregating ability (J. Amer. Chem. Soc., 115, 905-16 (1993). Substituted imidazolines were tested for their antidepressant activity in Ind. J.

- 6 Het. Chem., 2, 129-32 (1992). N-(4-Pyridy1) anthranilic amides were described in Chem Abstr. 97:109837 (1981). PCT publication W099/32477, published 1 July 1999, describes anthranilamides as anti-coagulants. US patent 6,140,351 describes anthranilamides as anti-coagulants. PCT
- publication W099/62885, published 9 December 1999, describes 1-(4-aminophenyl)pyrazoles as antiinflammatories. PCT publication W000/39111, published 6 July 2000, describes amides as factor Xa inhibitors. PCT publication W000/39117, published 6 July 2000, describes heteroaromatic amides as
- 15 factor Xa inhibitors. PCT publication W000/27819, published 18 May 2000, describes anthranilic acid amides as VEGF inhibitors. PCT publication W000/27820 published 18 May 2000, describes N-aryl anthranilic acid amides as VEGF inhibitors. 7-Chloroquinolinylamines are described in
- 20 FR2168227 as antiinflammatories. W001/55114, published 2 Aug. 2001, describes nicotinamides for the treatment of cancer. W001/55115, published 2 Aug. 2001, describes nicotinamides as inducers of apoptosis. W001/85715, published 15 November 2001, describes substituted pyridines
- 25 and pyrimidines as anti-angiogenesis agents. PCT publication W001/85691 published 15 November 2001, describes anthranilic amides as VEGF inhibitors. PCT publication W001/85671 published 15 November 2001, describes anthrany1 amides as VEGF inhibitors. PCT publication W001/81311
- 30 published 1 November 2001, describes anthranilic amides as VEGF inhibitors. However, compounds of the current invention have not been described as inhibitors of angiogenesis such as for the treatment of cancer.

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DESCRIPTION OF THE INVENTION

 $\,$ A class of compounds useful in treating cancer and $\,$ angiogenesis is defined by Formula I

$$R^2$$
 A
 A^2
 A^2
 A
 A^2

wherein each of A^1 and A^2 is independently C, CH or N; 10 wherein ring A is selected from

- a) 5- or 6-membered partially saturated heterocyclyl, preferably dihydropyran, dihydrothienyl, dihydrofuryl, oxo-dihydrofuryl, pyrrolinyl, dihydrothiazolyl, dihydro-oxazolyl, dihydro-isothiazolyl, dihydro-isoxazolyl, imidazolinyl and pyrazolinyl,
- b) 5- or 6-membered heteroaryl, preferably

selected from

I) 5-membered heteroaryl selected from thienyl, furanyl, pyrrolyl, thiazolyl, oxazolyl, imidazolyl, pyrazolyl, isoxazolyl, triazolyl and isothiazolyl, even more preferably 5-membered heteroaryl

5 A

$$R^{c}$$
 and R^{c} and R^{c}

II) preferably 6-membered heteroaryl selected from pyridyl, pyrazinyl, pyrimidinyl, pyridazinyl, and triazinyl,

even more preferably 6-membered heteroaryl selected from

more specifically

- c) 9-, 10- or 11-membered fused partially saturated heterocyclyl preferably tetrahydroquinolinyl,
- d) 9- or 10-membered fused heteroaryl, preferably
 - i) fused 9-membered fused heteroaryl selected from benzothienyl, benzothiazolyl, indolyl,

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benzimidazolyl, benzoxazolyl, benzofuryl, indazolyl and isoindolyl, and

- ii) fused 10-membered heteroaryl selected from quinolyl, isoquinolyl, naphthpyridinyl, quinoxalinyl and quinazolinyl,
- e) naphthyl, and
 - f) 4-, 5- or 6-membered cycloalkenyl, preferably 5-membered cycloalkenyl, more preferably cyclopentadienyl or cyclopentenyl;

wherein X is selected from

 \mathbb{Z} \mathbb{Z}

preferably X is selected from

N

more preferably X is
wherein Z is oxygen or sulfur;
wherein Y is selected from

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preferably Y is selected from

more preferably Y is selected from

even more preferably Y is -NH-CH2-;

wherein R^{a} and R^{b} are independently selected from H, halo,

cyano and C_{1-4} -alkyl substituted with R^2 , or wherein R^a and R^b together form C_3 - C_4 cycloalkyl,

preferably H, halo, cyano and C_{1-2} -alkyl substituted with R^2 , or wherein R^a and R^b together form C_1 - C_4 cycloalkyl, more preferably H, halo and C_1 - C_2 -alkyl,

even more preferably H;

15 wherein R^z is selected from C_1 - C_4 alkylenyl, where one of the CH_2 groups may be substituted with an oxygen atom or an - NH-.

preferably C1-C2 alkylenyl, where one of the CH2 groups
may be substituted with an oxygen atom or an -NHmore preferably C1-C2 alkylenyl;

wherein Rd is cycloalkyl,

preferably C₃-C₆ cycloalkyl; wherein R is selected from A-733A - 11
a) substituted or unsubstituted 5-6 membered heterocyclyl, preferably substituted or unsubstituted 5-6 membered heteroaryl comprising one or more nitrogen atoms, more preferably 4-pyrazolyl, triazolyl, 4-pyridyl, 4-pyrimidinyl, 5-pyrimidinyl, 6-pyrimidinyl, 4-pyridazinyl, or 6-pyridazinyl

pyridyl, 4-pyrimidinyl, 5-pyrimidinyl, 6pyrimidinyl, 4-pyridazinyl, or 6-pyridazinyl,
even more preferably 4-pyridyl, 4-pyrimidinyl
and 4-pyridazinyl,

10 even more preferably 4-pyridyl, and

b) substituted or unsubstituted fused 9-, 10- or 11membered heterocyclyl.

preferably substituted or unsubstituted 9-10 membered
fused heteroaryl comprising one or more nitrogen
atoms.

more preferably indazolyl, quinolinyl,
 isoquinolinyl, or quinazolinyl,
 even more preferably indazolyl, 4-quinolyl, 5 quinolyl, 6-quinolyl, 4-isoquinolyl, 5 isoquinolyl, and 6-isoquinolyl.

wherein substituted R is substituted with one or more substituents independently selected from halo, -OR³, -SR³, -SO₂R³, -CO₂R³, -CONR³R³, -COR³, -NR³R³, -SO₂NR³R³, -NR³C(0)OR³, -NR³C(0)R³, cycloalkyl, optionally substituted 5-6 membered heterocyclyl, optionally substituted phenyl, lower alkyl substituted with R², cyano, nitro, lower alkenyl and lower alkynyl; preferably halo, -OR³, -SR³, -CO₂R³, -CONR³R³, -SO₂NR³R³, -NR³C(0)OR³, -NR³C(0)R³, -NR³C(0)R³, cycloalkyl, optionally

substituted 5-6 membered heterocyclyl, optionally substituted phenyl, C₁₋₂-alkyl, cyano, C₁₋₂-hydroxyalkyl, nitro and C₁₋₂-haloalkyl;

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1.0

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wherein R1 is selected from

a) substituted or unsubstituted 6-10 membered aryl, preferably phenyl, naphthyl, indenyl, or tetrahydronaphthyl.

5 more preferably phenyl,

 b) substituted or unsubstituted 5-6 membered heterocyclyl,

preferably 5-6 membered heteroaryl,

more preferably thienyl, pyridyl, pyrimidinyl,
 pyridazinyl, pyrazolyl, imidazolyl, oxazolyl,
 thiazolyl, thiadiazolyl, furyl, or pyrrolyl,

 c) substituted or unsubstituted 9-10 membered fused heterocyclyl,

preferably 9-10 membered fused heteroaryl,

more preferably indazolyl, indolyl, 2,1,3benzothiadiazolyl, isoquinolyl, quinolyl,
tetrahydroquinolyl, benzodioxanyl, or
quinazolinyl,

- d) cycloalkyl, and
- e) cycloalkenyl

wherein substituted R¹ is substituted with one or more substituents independently selected from halo, -OR³, -SR³, -CO₂R³, -CONR³R³, -COR³, -NR³R³, -NH(C₁-C₄ alkylenylR¹⁴), -SO₂R³, -SO₂NR³R³, -NR³C(0)OR³, -NR³C(0)R³, optionally substituted cycloalkyl, optionally substituted 5-6 membered heterocyclyl, optionally substituted phenyl, lower alkyl substituted with R², cyano, nitro, lower alkenyl and lower alkynyl,

preferably R¹ is unsubstituted or substituted with
 one or more substituents independently selected
 from halo, -OR³, -SR³, -SO₂R³, -CO₂R³, -CONR³R³,
 -COR³, -NR³R³, -NH(C₁-C₂ alkylenylR³), -(C₁-C₂
 alkylenyl)NR³R³, -SO₂NR³R², -NR³C(O)OR³, -NR³C(O)R³,

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	optionally substituted cycloalkyl, optionally
	substituted 5-6 membered heterocyclyl, optionally
	substituted phenyl, optionally substituted
	phenyl- C_{1-2} -alkylenyl, optionally substituted 5-6
5	membered heterocyclyl- C_1 - C_2 -alkylenyl, C_{1-2} -alkyl,
	cyano, C_{1-2} -hydroxyalkyl, nitro and C_{1-2} -haloalkyl,
	more preferably R1 is unsubstituted or
	substituted with one or more substituents
	selected from chloro, fluoro, bromo, methoxy,
10	phenyloxy, benzyl, methylthio, methyl, ethyl,
	trifluoromethyl, difluoromethyl,
	pentafluoroethyl, hydroxymethyl, cyano,
	carboxy, aminocarbonyl, methylcarbonyl, amino,
	methylamino, cyclopropyl, cyclohexyl,
15	piperidinyl, morpholinyl, N-methylpiperazinyl,
	N-ethylpiperazinyl, morpholinylmethyl,
	methylpiperdinylmethyl,
	methylpiperazinylmethyl,
	methylaminothiocarbonyl, N-methylamino-
20	methylenyl, optionally substituted phenyl,
	N,N-diethylamino, or N,N-dimethylamino;
	wherein $\ensuremath{\mbox{R}^2}$ is one or more substituents independently selected
	from H, halo, $-OR^3$, oxo, $-SR^3$, $-CO_2R^3$, $-COR^3$, $-CONR^3R^3$,
	$-NR^3R^3,\ -SO_2NR^3R^3,\ -NR^3C(O)OR^3,\ -NR^3C(O)R^3,\ \text{cycloalkyl},$
25	optionally substituted phenylalkylenyl, optionally
	substituted 5-6 membered heterocyclyl, optionally
	substituted heteroarylalkylenyl, optionally substituted
	phenyl, lower alkyl, cyano, lower hydroxyalkyl, lower
	carboxyalkyl, nitro, lower alkenyl, lower alkynyl, lower
30	aminoalkyl, lower alkylaminoalkyl and lower haloalkyl,
	preferably \mathbb{R}^2 is one or more substituents independently
	selected from H, halo, $-OR^3$, oxo, $-SR^3$, $-CO_2R^3$,
	$-CONR^3R^3$, $-COR^3$, $-NR^3R^3$, $-SO_2NR^3R^3$, $-NR^3C$ (O)OR ³ ,
	$-NR^3C(0)R^3$, cycloalkyl, optionally substituted 5-6

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membered heterocyclyl, optionally substituted phenyl, C1-2-alkyl, cyano, C1-2-hydroxyalkyl, C1-3-carboxyalkyl, nitro, C2-3-alkenvl, C2-3-alkvnvl and C1-2-haloalkvl; wherein R3 is selected from H, lower alkyl, phenyl, 5-6 5 membered heterocyclyl, C1-C6 cycloalkyl, and lower haloalkvl. preferably H, C1-2-alkyl, phenyl, C3-C6 cycloalkyl, and C1--haloalkvl. more preferably H. methyl, phenyl, cyclopropyl, 10 cyclohexyl, and trifluoromethyl; wherein R4 is independently selected from C2-4-alkylenyl, C2-4-alkenylenyl and C2-4-alkynylenyl, where one of the CH2 groups may be substituted with an oxygen atom or an -NH-, preferably C2-3-alkylenyl where one of the CH2 groups 15 may be substituted with an oxygen atom or an -NH-, more preferably Co-Co alkylenyl: wherein R5 is selected from H, lower alkyl, phenyl and lower aralkyl, preferably H, methyl or ethyl; 20 wherein R6 is selected from H or C1.6-alkyl, preferably H or C1-2 alkyl; and wherein Rc is selected from H. methyl and optionally substituted phenyl; wherein R14 is selected from H. phenvl. 5-6 membered 25 heterocyclyl and C1-C6 cycloalkyl; wherein p is 0 to 2, preferably p is 2; and pharmaceutically acceptable salts thereof; provided A is not naphthyl when X is -C(O)NH- and when R1 is phenyl when Y is -NHCH2- and when R is 4-pyridyl; further 30 provided A is not pyridyl when X is -C(0)NH- and when R1 is 4-[3,5-bis(trifluoromethyl)-1H-pyrazol-1-yl]phenyl when Y is -N(CH3) - and when R is 4-methylpiperidinyl; further provided A is not pyridyl when X is -C(O)NH- and when Y is -NHCH2 and when R is 4-pyridylpiperidin-4-yl; 1-tertbutylpiperidin-

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4-yl, 1-isopropylpiperidin-4-yl or 1-cycloalkylpiperidin-4-yl; further provided A is not pyridyl when X is -C(0)NH- and when R¹ is 4-[3-(3-pyridyl)-5-(trifluoromethyl)-1H-pyrazol-1-yl]phenyl when Y is -NHCH₂- and when R is 4-pyridyl; and further provided R is not unsubstituted 2-thienyl, 2-pyridyl or 3-pyridyl.

The invention also relates to compounds of Formula II

II

wherein Ra and Rb are independently selected from H, halo,

 C_{1-4} -alkyl and $-N(R^6)_2$,

preferably H;

wherein n is 0-2;

preferably 1-2;

wherein R is selected from

- a) unsubstituted or substituted 5- or 6-membered nitrogen-containing heteroaryl, and
- b) unsubstituted or substituted 9- or 10-membered fused nitrogen-containing heteroaryl, preferably 4-pyridyl, pyrimidinyl, triazolyl,

pyridazinyl, indolyl, isoindolyl, indazolyl, quinolyl, isoquinolyl, naphthyridinyl or quinozalinyl.

where R is substituted with one or more substituents selected from halo, amino, hydroxy, C₁₋₆-alkyl, C₁₋₆-haloalkyl and C₁₋₆-alkoxy,

preferably substituted with one or more substituents selected from chloro, fluoro, amino, hydroxy, methyl, ethyl, propyl, trifluoromethyl, methoxy and ethoxy; A-733A - 16 -

wherein R1 is selected from unsubstituted or substituted arvl, 5-6-membered heteroarvl and 9-10 membered fused heteroarvl. preferably unsubstituted or substituted phenyl, 5 tetrahydronaphthyl, naphthyl, isoguinolyl, guinolyl, pyridyl, pyrimidinyl, pyridazinyl, indolyl, isoindolyl, naphthyridinyl, quinozalinyl, tetrahydroguinolinyl, indazolyl, benzothienyl, benzofuryl, benzimidazolyl, benzoxazolyl, or 10 benzthiazolvl. wherein R1 is substituted with one or more substituents selected from halo, C1-6-alkyl, optionally substituted C3-6-cycloalkyl, optionally substituted phenyl, C1-6-haloalkoxy, optionally substituted 15 phenyloxy, benzyl, optionally substituted 5-6 membered heterocyclyl-C, C,-alkylenyl, optionally substituted heteroarvl, optionally substituted heteroaryloxy, C1-6-haloalkyl, and C1-6-alkoxy, preferably chloro, fluoro, amino, hydroxy, 20 cyclohexyl, phenylmethyl, morpholinylmethyl, methylpiperdinylmethyl, methylpiperazinylmethyl, ethyl, propyl, trifluoromethyl, phenyloxy, methoxy and ethoxy; wherein R2 is one or more substituents independently 25 selected from Н. halo. C_{1-6} -alkyl, C1-6-haloalkyl, 30 C_{1-6} -alkoxy, C1-6-haloalkoxy, C1-6-carboxyalkyl, unsubstituted or substituted aryl and

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unsubstituted or substituted 5-6 membered heteroarvl:

preferably one or more substituents independently selected
 from H, chloro, fluoro, bromo, amino, hydroxy, methyl,
 ethyl, propyl, trifluoromethyl, methoxy, ethoxy,
 trifluoromethoxy, carboxymethyl, unsubstituted or
 substituted phenyl and unsubstituted or substituted
 heteroaryl selected

from thienyl, furanyl, pyridyl, imidazolyl, and
pyrazolyl; and

wherein R6 is H or C1-2-alkyl;

and pharmaceutically acceptable isomers and salts thereof.

The invention also relates to compounds of Formula III

III

15 wherein R^a and R^b are independently selected from H, halo, $C_{1-4}\text{-alkyl} \ \ \text{and} \ \ -N\left(R^6\right)_2,$

preferably H:

wherein n is 0-2;

preferably 1-2;

- 20 wherein R is selected from
 - a) unsubstituted or substituted 5- or 6-membered nitrogen-containing heteroaryl, and
 - b) unsubstituted or substituted 9- or 10-membered fused nitrogen-containing heteroaryl,
- 25 preferably 4-pyridyl, pyrimidinyl, pyridazinyl, indolyl, isoindolyl, indazolyl, quinolyl, isoquinolyl, naphthyridinyl or quinozalinyl,

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where R is substituted with one or more substituents
                 selected from halo, amino, hydroxy, C1-6-alkyl,
                 C1-6-haloalkyl and C1-6-alkoxy,
                preferably substituted with one or more
 5
                 substituents selected from chloro, fluoro,
                 amino, hydroxy, methyl, ethyl, propyl,
                 trifluoromethyl, methoxy and ethoxy:
     wherein R1 is selected from unsubstituted or substituted
       aryl, 5-6-membered heteroaryl and 9-10 membered fused
10
       heteroarvl.
       preferably unsubstituted or substituted phenyl,
          tetrahydronaphthyl, naphthyl, isoquinolyl, quinolyl,
          pyridyl, pyrimidinyl, pyridazinyl, indolyl,
          isoindolyl, naphthyridinyl, guinozalinyl,
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          tetrahydroguinolinyl, indazolyl, benzothienyl,
          benzofuryl, benzimidazolyl, benzoxazolyl, or
          benzthiazolyl,
          wherein R1 is substituted with one or more substituents
              selected from halo, C1_6-alkvl, optionally
20
              substituted C1-6-cycloalkyl, optionally substituted
              phenyl, C1.4-haloalkoxy, optionally substituted
              phenyloxy, benzyl, optionally substituted 5-6
              membered heterocyclyl-C1-C2-alkylenyl, optionally
              substituted heteroaryl, optionally substituted
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              heteroaryloxy, C1-6-haloalkyl, and C1-6-alkoxy,
             preferably chloro, fluoro, amino, hydroxy,
                cyclohexyl, phenylmethyl, morpholinylmethyl,
                methylpiperdinylmethyl, methylpiperazinylmethyl,
                ethyl, propyl, trifluoromethyl, phenyloxy,
3.0
                methoxy and ethoxy;
     wherein R2 is one or more substituents independently
          selected from
                Н.
                halo,
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C₁₋₆-alkyl,
C₁₋₆-haloalkyl,
C₁₋₆-alkoxy,
C₁₋₆-haloalkoxy,

C1_6-carboxvalkvl.

unsubstituted or substituted aryl and
unsubstituted or substituted 5-6 membered
heteroaryl;

preferably one or more substituents independently selected from H, chloro, fluoro, bromo, amino, hydroxy, methyl, ethyl, propyl, trifluoromethyl, methoxy, ethoxy, trifluoromethoxy, carboxymethyl, unsubstituted or substituted phenyl and unsubstituted or substituted heteroaryl selected from thienyl, furanyl, pyridyl, imidazolyl, and pyrazolyl; and

wherein R⁶ is H or C₁₋₂-alkyl;

and pharmaceutically acceptable isomers and salts thereof.

The invention also relates to compounds of Formula IV

$$\begin{array}{c|c} & O & \\ &$$

wherein A³ is selected from CR² and N;

wherein A^4 is selected from CR^2 and N; provided one of A^3 and A^4 is not CR^2 ;

ΙV

wherein Ra and Rb are independently selected from H, halo,

25 C_{1-4} -alkyl and -N(R⁶)₂,

preferably H;

wherein n is 0-2;

preferably 1-2;

wherein R is selected from

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	a) unsubstituted or substituted 5- or 6-membered
	nitrogen-containing heteroaryl, and
	b) unsubstituted or substituted 9- or 10-membered
	fused nitrogen-containing heteroaryl,
5	preferably 4-pyridyl, pyrimidinyl, pyridazinyl,
	indolyl, isoindolyl, indazolyl, quinolyl,
	isoquinolyl, naphthyridinyl or quinozalinyl,
	where R is substituted with one or more substituents
	selected from halo, amino, hydroxy, C_{1-6} -alkyl,
10	C_{1-6} -haloalkyl and C_{1-6} -alkoxy,
	preferably substituted with one or more
	substituents selected from chloro, fluoro,
	amino, hydroxy, methyl, ethyl, propyl,
	trifluoromethyl, methoxy and ethoxy;
15	wherein $\mathbf{R}^{\mathbf{l}}$ is selected from unsubstituted or substituted
	aryl, 5-6-membered heteroaryl and 9-10 membered fused
	heteroaryl,
	preferably unsubstituted or substituted phenyl,
	tetrahydronaphthyl, naphthyl, isoquinolyl, quinolyl,
20	pyridyl, pyrimidinyl, pyridazinyl, indolyl,
	isoindolyl, naphthyridinyl, quinozalinyl,
	tetrahydroquinolinyl, indazolyl, benzothienyl,
	benzofuryl, benzimidazolyl, benzoxazolyl, or
	benzthiazolyl,
25	wherein R1 is substituted with one or more substituents
	selected from halo, C_{1-6} -alkyl, optionally
	substituted C_{3-6} -cycloalkyl, optionally substituted
	phenyl, C_{1-6} -haloalkoxy, optionally substituted
	phenyloxy, benzyl, optionally substituted 5-6
30	membered heterocyclyl- C_1 - C_2 -alkylenyl, optionally
	substituted heteroaryl, optionally substituted
	heteroaryloxy, C_{1-6} -haloalkyl, and C_{1-6} -alkoxy,
	preferably chloro, fluoro, amino, hydroxy,

cyclohexyl, phenylmethyl, morpholinylmethyl,

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methylpiperdinylmethyl, methylpiperazinylmethyl,
ethyl, propyl, trifluoromethyl, phenyloxy,
methoxy and ethoxy;

wherein $\ensuremath{R^2}$ is one or more substituents independently

5 selected from

н.

halo,

 C_{1-6} -alkyl,

C₁₋₆-haloalkyl,

 C_{1-6} -alkoxy,

 C_{1-6} -haloalkoxy,

C₁₋₆-carboxyalkyl,

unsubstituted or substituted aryl and unsubstituted or substituted 5-6 membered

heteroarvl:

preferably one or more substituents independently selected from H, chloro, fluoro, bromo, amino, hydroxy, methyl, ethyl, propyl, trifluoromethyl, methoxy, ethoxy, trifluoromethoxy, carboxymethyl, unsubstituted or substituted phenyl and unsubstituted or substituted heteroaryl selected from thienyl, furanyl, pyridyl, imidazolyl, and pyrazolyl: and

wherein R⁶ is H or C₁₋₂-alkyl;

 $25\,$ $\,$ and pharmaceutically acceptable isomers and salts thereof.

The invention also relates to compounds of Formula V

wherein A5 is selected from S, O and NR6;

- 22 -

wherein Ra and Rb are independently selected from H. halo.

```
C_{1-4}-alkyl and -N(R^6)_2,
          preferably H:
     wherein n is 0-2;
 5
       preferably 1-2:
     wherein R is selected from
          a) unsubstituted or substituted 5- or 6-membered
               nitrogen-containing heteroarvl, and
          b) unsubstituted or substituted 9- or 10-membered
                fused nitrogen-containing heteroarvl.
10
              preferably 4-pyridyl, pyrimidinyl, pyridazinyl,
                 indolvl, isoindolvl, indazolvl, quinolvl,
                 isoguinolyl, naphthyridinyl or guinozalinyl,
            where R is substituted with one or more substituents
15
                 selected from halo, amino, hydroxy, C1.6-alkyl,
                 C1-6-haloalkyl and C1-6-alkoxy,
                preferably substituted with one or more
                 substituents selected from chloro, fluoro,
                 amino, hydroxy, methyl, ethyl, propvl.
                 trifluoromethyl, methoxy and ethoxy;
20
     wherein R1 is selected from unsubstituted or substituted
       aryl, 5-6-membered heteroaryl and 9-10 membered fused
       heteroarvl,
       preferably unsubstituted or substituted phenyl,
25
          tetrahydronaphthyl, naphthyl, isoquinolyl, quinolyl,
          pyridyl, pyrimidinyl, pyridazinyl, indolyl,
          isoindolyl, naphthyridinyl, quinozalinyl,
          tetrahydroquinolinyl, indazolyl, benzothienyl,
          benzofuryl, benzimidazolyl, benzoxazolyl, or
30
          benzthiazolvl.
          wherein R1 is substituted with one or more substituents
              selected from halo, C1-6-alkyl, optionally
              substituted C3-6-cycloalkyl, optionally substituted
              phenyl, C1-6-haloalkoxy, optionally substituted
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	phenyloxy, benzyl, optionally substituted 5-6
	membered heterocyclyl-C ₁ -C ₂ -alkylenyl, optionally
	substituted heteroaryl, optionally substituted
	heteroaryloxy, C_{1-6} -haloalkyl, and C_{1-6} -alkoxy,
5	preferably chloro, fluoro, amino, hydroxy,
	cyclohexyl, phenylmethyl, morpholinylmethyl,
	methylpiperdinylmethyl, methylpiperazinylmethyl,
	ethyl, propyl, trifluoromethyl, phenyloxy,
	methoxy and ethoxy;
10	wherein R ² is one or more substituents independently
	selected from
	Н,
	halo,
	C_{1-6} -alkyl,
15	C ₁₋₆ -haloalkyl,
	C_{1-6} -alkoxy,
	C ₁₋₆ -haloalkoxy,
	C ₁₋₆ -carboxyalkyl,
	unsubstituted or substituted aryl and
20	unsubstituted or substituted 5-6 membered
	heteroaryl;
	preferably one or more substituents independently
	selected from H, chloro, fluoro, bromo, amino,
	hydroxy, methyl, ethyl, propyl, trifluoromethyl,
25	methoxy, ethoxy, trifluoromethoxy, carboxymethyl,
	unsubstituted or substituted phenyl and
	unsubstituted or substituted heteroaryl selected
	from thienyl, furanyl, pyridyl, imidazolyl, and
	pyrazolyl; and
30	wherein R^6 is H or C_{1-2} -alkyl;
	and pharmaceutically acceptable isomers and salts thereof

The invention also relates to compounds of Formula VI

A-733A - 24 -

wherein A^5 is selected from S, O and NR 6 ; wherein R^a and R^b are independently selected from H, halo, $C_{1\text{-}4}\text{-alkyl} \text{ and } \text{-N}(R^6)_2,$

5 preferably H;

wherein n is 0-2;

preferably 1-2;

wherein R is selected from

 a) unsubstituted or substituted 5- or 6-membered nitrogen-containing heteroarvl, and

 b) unsubstituted or substituted 9- or 10-membered fused nitrogen-containing heteroaryl,
 preferably 4-pyridyl, pyrimidinyl, pyridazinyl,
 indolyl, isoindolyl, indazolyl, quinolyl,
 isoquinolyl, naphthyridinyl or quinozalinyl,

where R is substituted with one or more substituents selected from halo, amino, hydroxy, C₁₋₆-alkyl, C₁₋₆-haloalkyl and C₁₋₆-alkoxy,

preferably substituted with one or more substituents selected from chloro, fluoro, amino, hydroxy, methyl, ethyl, propyl, trifluoromethyl, methoxy and ethoxy;

wherein \mathbb{R}^1 is selected from unsubstituted or substituted aryl, 5-6-membered heteroaryl and 9-10 membered fused heteroaryl,

preferably unsubstituted or substituted phenyl,
 tetrahydronaphthyl, naphthyl, isoquinolyl, quinolyl,
 pyridyl, pyrimidinyl, pyridazinyl, indolyl,
 isoindolyl, naphthyridinyl, quinozalinyl,

TOTHEROT DITION

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A-733A - 25 -

tetrahydroguinolinyl, indazolyl, benzothienyl, benzofuryl, benzimidazolyl, benzoxazolyl, or benzthiazolvl. wherein R1 is substituted with one or more substituents 5 selected from halo, C1-6-alkyl, optionally substituted C1.6-cycloalkyl, optionally substituted phenyl, C1-6-haloalkoxy, optionally substituted phenyloxy, benzyl, optionally substituted 5-6 membered heterocyclyl-C1_C2-alkylenyl, optionally 10 substituted heteroarvl, optionally substituted heteroarvloxy, C1-6-haloalkyl, and C1-6-alkoxy, preferably chloro, fluoro, amino, hydroxy, cyclohexyl, phenylmethyl, morpholinylmethyl, methylpiperdinylmethyl, methylpiperazinylmethyl, 15 ethyl, propyl, trifluoromethyl, phenyloxy, methoxy and ethoxy: wherein R2 is one or more substituents independently selected from н. 20 halo. C1-6-alkyl, C₁₋₆-haloalkyl, C1_6-alkoxy, C1-6-haloalkoxy, 25 C1-6-carboxvalkvl. unsubstituted or substituted aryl and unsubstituted or substituted 5-6 membered heteroarvl; preferably one or more substituents independently 30 selected from H. chloro, fluoro, bromo, amino, hydroxy, methyl, ethyl, propyl, trifluoromethyl, methoxy, ethoxy, trifluoromethoxy, carboxymethyl, unsubstituted or substituted

phenyl and unsubstituted or substituted heteroaryl selected

from thienyl, furanyl, pyridyl, imidazolyl, and
pyrazolyl; and

VII

5 wherein R⁶ is H or C₁₋₂-alkyl;

and pharmaceutically acceptable isomers and salts thereof.

The invention also relates to compounds of Formula VII

$$R^2 = \underbrace{ \begin{bmatrix} 1 & 3 & 4 & 1 & 1 \\ 1 & 3 & 4 & 1$$

wherein A5 is selected from S, O and NR6;

10 wherein R^a and R^b are independently selected from H, halo, C_{1.4}-alkyl and -N(R⁶),

preferably H;

wherein n is 0-2;

preferably 1-2;

15 wherein R is selected from

- a) unsubstituted or substituted 5- or 6-membered nitrogen-containing heteroaryl, and
- b) unsubstituted or substituted 9- or 10-membered fused nitrogen-containing heteroaryl, preferably 4-pyridyl, pyrimidinyl, pyridazinyl,

indolyl, isoindolyl, indazolyl, quinolyl, isoquinolyl, naphthyridinyl or quinozalinyl,

where R is substituted with one or more substituents selected from halo, amino, hydroxy, C1.s-alkyl,

 C_{1-6} -haloalkyl and C_{1-6} -alkoxy,

preferably substituted with one or more substituents selected from chloro, fluoro, amino, hydroxy, methyl, ethyl, propyl, trifluoromethyl, methoxy and ethoxy; A-733A - 27 -

wherein R1 is selected from unsubstituted or substituted arvl, 5-6-membered heteroarvl and 9-10 membered fused heteroarvl. preferably unsubstituted or substituted phenyl, 5 tetrahydronaphthyl, naphthyl, isoguinolyl, guinolyl, pyridyl, pyrimidinyl, pyridazinyl, indolyl, isoindolyl, naphthyridinyl, quinozalinyl, tetrahydroguinolinyl, indazolyl, benzothienyl, benzofuryl, benzimidazolyl, benzoxazolyl, or 10 benzthiazolvl. wherein R1 is substituted with one or more substituents selected from halo, C1-6-alkyl, optionally substituted C2-6-cvcloalkyl, optionally substituted phenyl, C1-6-haloalkoxy, optionally substituted 15 phenyloxy, benzyl, optionally substituted 5-6 membered heterocyclyl-C1-C2-alkylenyl, optionally substituted heteroaryl, optionally substituted heteroaryloxy, C1-6-haloalkyl, and C1-6-alkoxy, preferably chloro, fluoro, amino, hydroxy, 20 cyclohexyl, phenylmethyl, morpholinylmethyl, methylpiperdinylmethyl, methylpiperazinylmethyl, ethyl, propyl, trifluoromethyl, phenyloxy, methoxy and ethoxy: wherein R2 is one or more substituents independently 25 selected from н. halo, C_{1-6} -alkvl, C1-6-haloalkyl, 30 C1-6-alkoxy, C1-6-haloalkoxy, C1-6-carboxyalkyl, unsubstituted or substituted arvl and

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unsubstituted or substituted 5-6 membered heteroarvl;

preferably one or more substituents independently selected from H, chloro, fluoro, bromo, amino, hydroxy, methyl, ethyl, propyl, trifluoromethyl, methoxy, ethoxy, trifluoromethoxy, carboxymethyl, unsubstituted or substituted phenvl and unsubstituted or substituted heteroarvl selected from thienyl, furanyl, pyridyl, imidazolyl, and

pvrazolv1: and wherein R6 is H or C1-2-alkvl;

and pharmaceutically acceptable isomers and salts thereof.

The invention also relates to compounds of Formula

VIII

wherein A5 is selected from S, O and NR6; wherein Ra and Rb are independently selected from H, halo, C_{1-4} -alkyl and $-N(R^6)_2$.

preferably H;

20 wherein n is 0-2:

preferably 1-2;

wherein R is selected from

- a) unsubstituted or substituted 5- or 6-membered nitrogen-containing heteroarvl, and
- b) unsubstituted or substituted 9- or 10-membered 25 fused nitrogen-containing heteroaryl, preferably 4-pyridyl, pyrimidinyl, pyridazinyl, indolyl, isoindolyl, indazolyl, quinolyl, isoquinolyl, naphthyridinyl or quinozalinyl,

A-733A - 29 -

where R is substituted with one or more substituents selected from halo, amino, hydroxy, C1-6-alkyl, C1-6-haloalkyl and C1-6-alkoxy. preferably substituted with one or more 5 substituents selected from chloro, fluoro, amino, hydroxy, methyl, ethyl, propyl, trifluoromethyl, methoxy and ethoxy; wherein R1 is selected from unsubstituted or substituted arvl. 5-6-membered heteroarvl and 9-10 membered fused 10 heteroarvl. preferably unsubstituted or substituted phenyl. tetrahydronaphthyl, naphthyl, isoguinolyl, guinolyl, pyridyl, pyrimidinyl, pyridazinyl, indolyl, isoindolyl, naphthyridinyl, quinozalinyl, 15 tetrahydroguinolinyl, indazolyl, benzothienyl, benzofuryl, benzimidazolyl, benzoxazolyl, or benzthiazolvl. wherein R1 is substituted with one or more substituents selected from halo, C1-6-alkyl, optionally 20 substituted C1.6-cvcloalkvl, optionally substituted phenyl, C1-6-haloalkoxy, optionally substituted phenyloxy, benzyl, optionally substituted 5-6 membered heterocyclyl-C1-C2-alkylenyl, optionally substituted heteroaryl, optionally substituted 25 heteroaryloxy, C1-6-haloalkyl, and C1-6-alkoxy, preferably chloro, fluoro, amino, hydroxy, cyclohexyl, phenylmethyl, morpholinylmethyl, methylpiperdinylmethyl, methylpiperazinylmethyl, ethyl, propyl, trifluoromethyl, phenyloxy, 3.0 methoxy and ethoxy: wherein R2 is one or more substituents independently selected from

> H, halo,

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C₁₋₆-alkyl,

C1-6-haloalkyl,

C1_6-alkoxy,

C1-6-haloalkoxy,

C₁₋₆-carboxyalkyl,

unsubstituted or substituted aryl and unsubstituted or substituted 5-6 membered

heteroaryl;

pyrazolyl; and

preferably one or more substituents independently selected from H, chloro, fluoro, bromo, amino, hydroxy, methyl, ethyl, propyl, trifluoromethyl, methoxy, ethoxy, trifluoromethoxy, carboxymethyl, unsubstituted or substituted phenyl and unsubstituted or substituted heteroaryl selected from thienyl, furanyl, pyridyl, imidazolyl, and

wherein R6 is H or C1-2-alkyl;

and pharmaceutically acceptable isomers and salts thereof.

The invention also relates to compounds of Formula IX

IX

wherein A⁵ is selected from S, O and NR⁶;

wherein \boldsymbol{R}^{a} and \boldsymbol{R}^{b} are independently selected from H, halo,

 C_{1-4} -alkyl and $-N(R^6)_2$,

preferably H;

25 wherein n is 0-2;

preferably 1-2;

wherein R is selected from

 a) unsubstituted or substituted 5- or 6-membered nitrogen-containing heteroaryl, and A-733A - 31 -

b) unsubstituted or substituted 9- or 10-membered

	fused nitrogen-containing heteroaryl,
	preferably 4-pyridyl, pyrimidinyl, pyridazinyl,
	indolyl, isoindolyl, indazolyl, quinolyl,
5	isoquinolyl, naphthyridinyl or quinozalinyl,
	where R is substituted with one or more substituents
	selected from halo, amino, hydroxy, C_{1-6} -alkyl,
	C_{1-6} -haloalkyl and C_{1-6} -alkoxy,
	preferably substituted with one or more
10	substituents selected from chloro, fluoro,
	amino, hydroxy, methyl, ethyl, propyl,
	trifluoromethyl, methoxy and ethoxy;
	wherein \mathbb{R}^1 is selected from unsubstituted or substituted
	aryl, 5-6-membered heteroaryl and 9-10 membered fused
15	heteroaryl,
	preferably unsubstituted or substituted phenyl,
	tetrahydronaphthyl, naphthyl, isoquinolyl, quinolyl,
	pyridyl, pyrimidinyl, pyridazinyl, indolyl,
	isoindolyl, naphthyridinyl, quinozalinyl,
20	tetrahydroquinolinyl, indazolyl, benzothienyl,
	benzofuryl, benzimidazolyl, benzoxazolyl, or
	benzthiazolyl,
	wherein R^1 is substituted with one or more substituent
	selected from halo, C_{1-6} -alkyl, optionally
25	substituted C_{3-6} -cycloalkyl, optionally substituted
	phenyl, C_{1-6} -haloalkoxy, optionally substituted
	phenyloxy, benzyl, optionally substituted 5-6
	membered heterocyclyl- C_1 - C_2 -alkylenyl, optionally
	substituted heteroaryl, optionally substituted
30	heteroaryloxy, C_{1-6} -haloalkyl, and C_{1-6} -alkoxy,
	preferably chloro, fluoro, amino, hydroxy,
	cyclohexyl, phenylmethyl, morpholinylmethyl,
	methylpiperdinylmethyl, methylpiperazinylmethyl,

ethyl, propyl, trifluoromethyl, phenyloxy, methoxy and ethoxy;

wherein ${\bf R}^2$ is one or more substituents independently selected from

5 н,

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halo.

C1_6-alkvl,

C1-6-haloalkyl,

 C_{1-6} -alkoxy,

C1-6-haloalkoxy,

C1-6-carboxyalkyl,

unsubstituted or substituted aryl and

unsubstituted or substituted 5-6 membered

heteroaryl;

preferably one or more substituents independently selected from H, chloro, fluoro, bromo, amino, hydroxy, methyl, ethyl, propyl,

trifluoromethyl, methoxy, ethoxy,

trifluoromethoxy, carboxymethyl, unsubstituted or substituted phenyl and unsubstituted or substituted heteroaryl selected

from thienyl, furanyl, pyridyl, imidazolyl, and
pyrazolyl; and

wherein R6 is H or C1-2-alkyl;

25 and pharmaceutically acceptable isomers and salts thereof.

The invention also relates to compounds of Formula X

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wherein A5 is selected from S, O and NR6;
     wherein A<sup>6</sup> is selected from N and CR<sup>2</sup>;
     wherein Ra and Rb are independently selected from H, halo,
          C_{1-4}-alkyl and -N(R^6)_2,
 5
          preferably H:
     wherein n is 0-2;
       preferably 1-2;
     wherein R is selected from
          a) unsubstituted or substituted 5- or 6-membered
10
               nitrogen-containing heteroaryl, and
          b) unsubstituted or substituted 9- or 10-membered
                fused nitrogen-containing heteroaryl,
               preferably 4-pyridyl, pyrimidinyl, pyridazinyl,
                  indolyl, isoindolyl, indazolyl, quinolyl,
15
                  isoquinolyl, naphthyridinyl or quinozalinyl,
            where R is substituted with one or more substituents
                 selected from halo, amino, hydroxy, C1-6-alkyl,
                 C1-6-haloalkyl and C1-6-alkoxy,
                preferably substituted with one or more
20
                  substituents selected from chloro, fluoro,
                 amino, hydroxy, methyl, ethyl, propyl,
                 trifluoromethyl, methoxy and ethoxy;
     wherein R1 is selected from unsubstituted or substituted
        aryl, 5-6-membered heteroaryl and 9-10 membered fused
25
       heteroaryl,
       preferably unsubstituted or substituted phenyl,
          tetrahydronaphthyl, naphthyl, isoguinolyl, guinolyl,
          pyridyl, pyrimidinyl, pyridazinyl, indolyl,
          isoindolyl, naphthyridinyl, quinozalinyl,
30
          tetrahydroguinolinyl, indazolyl, benzothienyl,
          benzofuryl, benzimidazolyl, benzoxazolyl, or
          benzthiazolyl,
          wherein R1 is substituted with one or more substituents
              selected from halo, C1-6-alkyl, optionally
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substituted C3-6-cycloalkyl, optionally substituted phenyl, C1-6-haloalkoxy, optionally substituted phenyloxy, benzyl, optionally substituted 5-6 membered heterocyclyl-C1-C2-alkylenyl, optionally substituted heteroaryl, optionally substituted heteroarvloxy, C1-6-haloalkyl, and C1-6-alkoxy, preferably chloro, fluoro, amino, hydroxy, cyclohexyl, phenylmethyl, morpholinylmethyl, methylpiperdinylmethyl, methylpiperazinylmethyl, ethyl, propyl, trifluoromethyl, phenyloxy, methoxy and ethoxy; wherein R2 is one or more substituents independently selected from н. halo. C1-6-alkyl, C1-6-haloalkyl, C_{1-6} -alkoxy, C1-6-haloalkoxy, C1-6-carboxyalkyl, unsubstituted or substituted arvl and unsubstituted or substituted 5-6 membered heteroaryl; preferably one or more substituents independently selected from H, chloro, fluoro, bromo, amino, hydroxy, methyl, ethyl, propyl, trifluoromethyl, methoxy, ethoxy, trifluoromethoxy, carboxymethyl, unsubstituted or substituted phenyl and unsubstituted or substituted heteroaryl selected from thienvl, furanvl, pyridyl, imidazolyl, and

wherein

pyrazolyl;

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wherein R⁶ is H or C₁₋₂-alkyl;

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and pharmaceutically acceptable isomers and salts thereof.

The invention also relates to compounds of Formula II'

II'

5 wherein R is selected from

 a) unsubstituted or substituted 5- or 6-membered nitrogen-containing heteroaryl,

preferably 4-pyridyl, 3-pyridyl, 2-pyridyl,
 pyrimidinyl, triazolyl, and pyridazinyl,
 more preferably 4-pyridyl, and

b) unsubstituted or substituted 9- or 10-membered fused heterocyclyl

preferably indolyl, isoindolyl, indazolyl, quinolyl, isoquinolyl, benzotriazolyl, 2,3- dihydrobenzofuryl, 2-oxo-1,2-dihydroquinol-7-yl, naphthyridinyl and quinozalinyl.

where substituted R is substituted with one or more

substituents selected from halo, amino, hydroxy, oxo, C₁₋₆-alkyl, C₁₋₆-haloalkyl, C₁₋₆-alkoxy, optionally substituted heterocyclyl-C₁₋₆-alkoxy, optionally substituted heterocyclyl-C₁₋₆-alkylamino, optionally substituted heterocyclyl-C₁₋₆-alkyl, C₁₋₆-alkylamino-C₂₋₄-alkynyl, C₁₋₆-alkylamino-C₁₋₆-alkoxy, C₁₋₆-alkylamino-C₁₋₆-alkoxy-C₁₋₆-alkoxy, and optionally substituted

heterocyclyl-C₂₋₄-alkynyl,
preferably chloro, fluoro, amino, hydroxy, methyl,
ethyl, propyl, trifluoromethyl,
dimethylaminopropynyl, 1-

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methylpiperdinylmethoxy,

dimethylaminoethoxyethoxy, methoxy and ethoxy; wherein R¹ is selected from unsubstituted or substituted aryl, preferably phenyl, tetrahydronaphthyl, indanyl, indenyl, and naphthyl.

cycloalkyl, preferably cyclohexyl,

5-6 membered heteroaryl, preferably isoxazolyl, pyrazolyl, thiazolyl, thiadiazolyl, thienyl, pyridyl, pyrimidinyl, and pyridazinyl, and

pyrimidinyi, and pyridazinyi, and

9-10 membered bicyclic and 13-14 membered tricyclic heterocyclyl, preferably 1,2-dihydroquinolyl, 1,2,3,4-tetrahydro-isoquinolyl, isoquinolyl, quinolyl, indolyl, isoindolyl, 2,3-dihydro-1H-indolyl, naphthyridinyl, quinozalinyl, benzo[d]isothiazolyl, 2,3,4,4a,9,9a-hexahydro-1H-3-aza-fluorenyl, 5,6,7-trihydro-1,2,4-triazolo[3,4-a]isoquinolyl, tetrahydroquinolinyl, indazolyl, 2,1,3-benzothiadiazolyl, benzodioxanyl,

benzothienyl, benzofuryl, dihydro-benzimidazolyl, benzimidazolyl, benzoxazolyl and benzthiazolyl;

wherein substituted R¹ is substituted with one or more substituents selected from halo, C₁₋₆-alkyl, optionally substituted C₂₋₆-cycloalkyl, optionally substituted phenyl, optionally substituted phenyl-C₁.C₄-alkylenyl, C₁₋₂-haloalkoxy, optionally substituted 4-6 membered heterocyclyl-C₁.C₄-alkylenyl, optionally substituted 4-6 membered heterocyclyl-C₂.C₄-alkenylenyl, optionally substituted 4-6 membered heterocyclyl, optionally substituted 4-6 membered heterocyclyl, optionally substituted phenyloxy, optionally substituted 4-6

membered heterocyclyloxy, optionally substituted 4-6 membered heterocyclyl-C₁₋₄-alkyloxy, optionally substituted 4-6 membered heterocyclylsulfonyl, optionally substituted 4-6 membered heterocyclylamino, optionally substituted 4-6 membered heterocyclylcarbonyl, optionally

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substituted 4-6 membered heterocyclyl- C_{1-4} -alkylcarbonyl, C_{1-2} -haloalkyl, C_{1-4} -aminoalkyl, nitro, amino, -NHC(0)NH₂, alkylcarbonylamino, hydroxy, oxo, cyano, aminosulfonyl, C_{1-2} -alkylsulfonyl, halosulfonyl, C_{1-4} -alkylcarbonyl, C_{1-3} -alkylamino- C_{1-3} -alkyl, C_{1-3} -alkylamino- C_{1-3} -alkoxy, C_{1-3} -alkylamino- C_{1-3} -alkoxy- C_{1-3} -alkoxy, C_{1-4} -alkoxycarbonyl, C_{1-4} -alkoxycarbonylamino- C_{1-4} -alkoxycarbonylamino- C_{1-4} -alkyl, C_{1-4} -hydroxyalkyl,

and C1-4-alkoxy,

preferably bromo, chloro, fluoro, iodo, nitro, amino, cyano, aminoethyl, Boc-aminoethyl. hydroxy, oxo, aminosulfonyl, 4methylpiperazinylsulfonyl, cyclohexyl, phenyl, phenylmethyl, morpholinylmethyl, 1methylpiperazin-4-vlmethyl, 1-methylpiperazin-4ylpropyl, morpholinylpropyl, piperidin-1ylmethyl, 1-methylpiperidin-4-ylmethyl, 2-methyl-2-(1-methylpiperidin-4-yl)ethyl, morpholinvlethyl, 1-(4-morpholinvl)-2,2dimethylpropyl, piperidin-4-ylethyl, 1-Bocpiperidin-4-ylethyl, piperidin-1-ylethyl, 1-Bocpiperidin-4-ylethyl, piperidin-4-ylmethyl, 1-Bocpiperidin-4-ylmethyl, piperidin-4-ylpropyl, 1-Boc-piperidin-4-vlpropvl, piperidin-1-vlpropvl, pyrrolidin-1-vlpropyl, pyrrolidin-2-vlpropyl, 1-Boc-pyrrolidin-2-ylpropyl, pyrrolidin-1-ylmethyl, pyrrolidin-2-ylmethyl, 1-Boc-pyrrolidin-2ylmethyl, pyrrolidinylpropenyl, pyrrolidinylbutenyl, fluorosulfonyl, methylsulfonyl, methylcarbonyl, Boc, piperidin-1ylmethylcarbonyl, 4-methylpiperazin-1ylcarbonylethyl, methoxycarbonyl, aminomethylcarbonyl, dimethylaminomethylcarbonyl,

3-ethoxycarbonyl-2-methyl-fur-5-yl, 4-

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methylpiperazin-1-vl, 4-methyl-1-piperidyl, 1-
                Boc-4-piperidyl, piperidin-4-yl, 1-
                methylpiperidin-4-vl, 1-methyl-(1,2,3,6-
                tetrahydropyridyl), imidazolyl, morpholinyl, 4-
 5
                trifluoromethyl-1-piperidinyl, hydroxybutyl,
                methyl, ethyl, propyl, isopropyl, butyl, tert-
                butyl, sec-butyl, trifluoromethyl,
                pentafluoroethyl, nonafluorobutyl,
                dimethylaminopropyl, 1,1-di(trifluoromethyl)-1-
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                hydroxymethyl, 1.1-di(trifluoromethyl)-1-
                (piperidinylethoxy)methyl, 1,1-
                di(trifluoromethyl)-1-
                (methoxyethoxyethoxy)methyl, 1-hydroxyethyl, 2-
                hydroxyethyl, trifluoromethoxy, 1-aminoethyl, 2-
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                aminoethyl, 1-(N-isopropylamino)ethyl, 2-(N-
                isopropylamino)ethyl, dimethylaminoethoxy, 4-
                chlorophenoxy, phenyloxy, azetidin-3-vlmethoxy,
                1-Boc-azetidin-3-ylmethoxy, pyrrol-2-ylmethoxy,
                1-Boc-pyrrol-2-ylmethoxy, pyrrol-1-ylmethoxy, 1-
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                methyl-pyrrol-2-vlmethoxy, 1-isopropyl-pyrrol-2-
                ylmethoxy, 1-Boc-piperdin-4-ylmethoxy, piperdin-
                4-vlmethoxy, 1-methylpiperdin-4-vloxy,
                isopropoxy, methoxy and ethoxy;
    wherein R2 is one or more substituents independently
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          selected from
                н.
                halo,
                hydroxy,
                amino.
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                C1-6-alkvl.
                C1-6-haloalkyl,
                C_{1-6}-alkoxy,
                C1-2-alkylamino,
                aminosulfonyl,
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C₃₋₆-cycloalkyl, cyano, C₁₋₂-hydroxyalkyl, nitro,

 C_{2-3} -alkenyl,

 C_{2-3} -alkynyl,

 C_{1-6} -haloalkoxy,

 C_{1-6} -carboxyalkyl,

5-6-membered heterocyclyl-C₁₋₆-alkylamino, unsubstituted or substituted phenyl and unsubstituted or substituted 5-6 membered heterocyclyl;

preferably H, chloro, fluoro, bromo, amino, hydroxy,
 methyl, ethyl, propyl, oxo, dimethylamino,
 aminosulfonyl, cyclopropyl, cyano, hydroxymethyl,
 nitro, propenyl, trifluoromethyl, methoxy, ethoxy,
 trifluoromethoxy, carboxymethyl,
 morpholinylethylamino, propynyl, unsubstituted or
 substituted phenyl and unsubstituted or substituted
 heteroaryl selected from thienyl,

furanyl, pyridyl, imidazolyl, and pyrazolyl; wherein R^4 is selected from a direct bond, C_{1-4} -alkyl, and

preferably a direct bond, ethyl, butyl, and wherein R² is selected from C₁₋₂-alkyl, C₂₋₆-branched alkyl, C₂₋₄-branched haloalkyl, amino-C₁₋₄-alkyl and C₁₋₂-alkylamino-C₁₋₂-alkyl,

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wherein Re and Rf are independently selected from H and C1-2haloalkvl. preferably trifluoromethyl; and wherein R7 is selected from H, C1-3-alkyl, optionally substituted phenyl, optionally substituted phenyl-C1-1alkyl, optionally substituted 4-6 membered heterocyclyl, optionally substituted 4-6 membered heterocyclyl-C1-C1-alkyl, C1-1-alkylamino-C1-1-alkyl, C1-1alkoxy-C1-2-alkyl and C1-3-alkoxy-C1-3-alkoxy-C1-3-alkyl; provided R2 is not H, or provided R1 is not heteroaryl or arvl or provided R is substituted with optionally substituted heterocyclyl-C1-6-alkoxy, optionally substituted heterocyclyl-C, a-alkylamino, optionally substituted heterocyclyl-C1-6-alkyl, C1-6-alkylamino-C2-4-alkynyl, C1-6-alkylamino-C1-6-alkoxy, C1-6-alkylamino-C1-6-alkoxy-C1-6-alkoxy, or optionally substituted heterocyclyl-C2-4-alkynyl, or R1 is substituted with optionally substituted phenyloxy, optionally substituted 5-6 membered heterocyclyloxy, optionally substituted 5-6 membered heterocyclylsulfonyl, optionally substituted 5-6 membered heterocyclylamino.

membered heterocyclyl-C₁₋₄-alkylcarbonyl, C₁₋₃25 alkylamino-C₁₋₃-alkoxy, or C₁₋₃-alkylamino-C₁₋₃-alkoxy-C₁₋₃
3-alkoxy; further provided R is not 3-pyridyl when R²
is CH₂;

heterocyclylcarbonyl, optionally substituted 5-6

optionally substituted 5-6 membered

and pharmaceutically acceptable isomers and derivatives thereof.

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The invention also relates to compounds of Formula XI

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wherein R is selected from

a) unsubstituted or substituted 5- or 6-membered nitrogen-containing heteroaryl, preferably 4-pyridyl, 3-pyridyl, 2-pyridyl, pyrimidinyl, triazolyl, and pyridazinyl,

more preferably 4-pyridyl, and

b) unsubstituted or substituted 9- or 10-membered fused heteroarvl

preferably indolyl, isoindolyl, indazolyl, quinolyl, isoquinolyl, benzotriazolyl, naphthyridinyl and quinozalinyl,

where substituted R is substituted with one or more substituents selected from halo, amino, hydroxy, \$C_{1-6}\$-alkyl, \$C_{1-6}\$-haloalkyl, \$C_{1-6}\$-alkoxy, optionally substituted heterocyclyl-\$C_{1-6}\$-alkylamino, optionally substituted heterocyclyl-\$C_{1-6}\$-alkylamino, optionally substituted heterocyclyl-\$C_{1-6}\$-alkylamino-\$C_{1-6}\$-alkylamino-\$C_{2-4}\$-alkynyl, \$C_{1-6}\$-alkylamino-\$C_{1-6}\$-alkoxy, \$C_{1-6}\$-alkoxy, and optionally substituted heterocyclyl-\$C_{2-4}\$-alkoxy, and optionally substituted heterocyclyl-\$C_{2-4}\$-alkynyl, preferably chloro, fluoro, amino, hydroxy, methyl, ethyl, propyl, trifluoromethyl,

dimethylaminopropynyl, 1-methylpiperdinylmethoxy.

 $\label{eq:dimethylaminoethoxyethoxy, methoxy} \mbox{ and ethoxy;} \\ \mbox{wherein } R^1 \mbox{ is selected from unsubstituted or substituted} \\ \mbox{ arvl.}$

cycloalkyl, 5-6 membered heteroarvl and 9-10 membered bicyclic and 13-14 membered tricyclic heterocyclyl, preferably phenyl, tetrahydronaphthyl, indanyl, 5 indenvl, naphthyl, cyclohexyl, isoxazolyl, pyrazolyl, thiazolyl, thiadiazolyl, thienyl, pyridyl, pyrimidinyl, pyridazinyl, 1,2dihydroquinolyl, 1,2,3,4-tetrahydro-isoquinolyl, 10 isoquinolyl, quinolyl, indolyl, isoindolyl, 2,3dihydro-1H-indolyl, naphthyridinyl, quinozalinyl, benzo[dlisothiazolvl, 2.3.4.4a.9.9a-hexahvdro-1H-3aza-fluorenvl, 5,6,7-trihvdro-1,2,4-triazolo[3,4alisoguinolyl, tetrahydroguinolinyl, indazolyl, 15 2,1,3-benzothiadiazolyl, benzodioxanyl, benzothienvl, benzofurvl, dihydro-benzimidazolyl, benzimidazolyl, benzoxazolyl and benzthiazolyl, specifically 4-6 membered saturated or partially un-saturated monocyclic heterocyclyl, 20 9-10 membered saturated or partially unsaturated bicyclic heterocyclyl, and 13-14 membered saturated or partially unsaturated tricyclic heterocyclyl, more specifically 1,2-dihydroquinolyl, 25 1,2,3,4-tetrahydro-isoguinolyl, 2,3-dihydro-1H-indolyl, benzo[d]isothiazolyl, dihydrobenzimidazolyl, 2,3,4,4a,9,9a-hexahydro-1H-3aza-fluorenvl, 5,6,7-trihvdro-1,2,4-

wherein substituted R¹ is substituted with one or more substituents selected from halo, C₁₋₆-alkyl, optionally substituted C₁₋₆-cycloalkyl, optionally substituted phenyl, optionally substituted phenyl-C_{1-C4}-alkylenyl,

tetrahydroquinolinyl,

triazolo[3,4-a]isoquinolyl, and

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C1-2-haloalkoxy, optionally substituted 4-6 membered heterocyclyl-C1-C4-alkyl, optionally substituted 4-6 membered heterocyclyl-C2-C4-alkenyl, optionally substituted 4-6 membered heterocyclyl, optionally substituted phenyloxy, optionally substituted 4-6 membered heterocyclyloxy, optionally substituted 4-6 membered heterocyclyl-C1-C4-alkoxy, optionally substituted 4-6 membered heterocyclylsulfonyl, optionally substituted 4-6 membered heterocyclylamino, optionally substituted 4-6 membered heterocyclylcarbonyl, optionally substituted 5-6 membered heterocyclyl-C1_4-alkylcarbonyl, C1_2haloalkyl, C1-4-aminoalkyl, nitro, amino, hydroxy, oxo, cyano, aminosulfonyl, C1-2-alkylsulfonyl, halosulfonyl, C1_4-alkylcarbonyl, C1_3-alkylamino-C1_3-alkyl, C1_3alkylamino-C1-3-alkoxy, C1-3-alkylamino-C1-3-alkoxy-C1-3alkoxy, C1-4-alkoxycarbonyl, C1-4-alkoxycarbonylamino-C1-

4-alkvl, C1-4-hvdroxvalkvl, and C1_4-alkoxy, preferably bromo, chloro, fluoro, iodo, nitro, amino, cyano, aminoethyl, Boc-aminoethyl, hydroxy, oxo, aminosulfonvl, 4-methylpiperazinvlsulfonvl, cyclohexyl, phenyl, phenylmethyl, morpholinylmethyl, 1-methylpiperazin-4-ylmethyl, 1-methylpiperazin-4-ylpropyl, morpholinylpropyl, piperidin-1-vlmethyl, 1-methylpiperidin-4ylmethyl, 2-methyl-2-(1-methylpiperidin-4yl)ethyl, morpholinylethyl, 1-(4-morpholinyl)-2,2-dimethylpropyl, piperidin-4-ylethyl, 1-Bocpiperidin-4-ylethyl, piperidin-1-ylethyl, 1-Bocpiperidin-4-ylethyl, piperidin-4-ylmethyl, 1-Bocpiperidin-4-vlmethyl, piperidin-4-vlpropyl, 1-Boc-piperidin-4-ylpropyl, piperidin-1-ylpropyl,

pyrrolidin-1-vlpropyl, pyrrolidin-2-vlpropyl, 1-

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		Boc-pyrrolidin-2-ylpropyl, pyrrolidin-1-ylmethyl,
		pyrrolidin-2-ylmethyl, 1-Boc-pyrrolidin-2-
		ylmethyl, pyrrolidinylpropenyl,
		pyrrolidinylbutenyl, fluorosulfonyl,
5	:	methylsulfonyl, methylcarbonyl, Boc, piperidin-1-
	:	ylmethylcarbonyl, 4-methylpiperazin-1-
		ylcarbonylethyl, methoxycarbonyl,
		aminomethylcarbonyl, dimethylaminomethylcarbonyl,
		3-ethoxycarbonyl-2-methyl-fur-5-yl, 4-
10	:	methylpiperazin-1-yl, 4-methyl-1-piperidyl, 1-
		Boc-4-piperidyl, piperidin-4-yl, 1-
	;	methylpiperidin-4-yl, 1-methyl-(1,2,3,6-
		tetrahydropyridyl), imidazolyl, morpholinyl, 4-
		trifluoromethyl-1-piperidinyl, hydroxybutyl,
15	:	methyl, ethyl, propyl, isopropyl, butyl, tert-
	;	butyl, sec-butyl, trifluoromethyl,
		pentafluoroethyl, nonafluorobutyl,
		dimethylaminopropyl, 1,1-di(trifluoromethyl)-1-
	:	hydroxymethyl, 1,1-di(trifluoromethyl)-1-
20		(piperidinylethoxy)methyl, 1,1-
		di(trifluoromethyl)-1-
		(methoxyethoxyethoxy) methyl, 1-hydroxyethyl, 2-
	:	hydroxyethyl, trifluoromethoxy, 1-aminoethyl, 2-
		aminoethyl, 1-(N-isopropylamino)ethyl, 2-(N-
25		isopropylamino)ethyl, dimethylaminoethoxy, 4-
		chlorophenoxy, phenyloxy, azetidin-3-ylmethoxy,
		1-Boc-azetidin-3-ylmethoxy, pyrrol-2-ylmethoxy,
		1-Boc-pyrrol-2-ylmethoxy, pyrrol-1-ylmethoxy, 1-
		methyl-pyrrol-2-ylmethoxy, 1-isopropyl-pyrrol-2-
30	:	ylmethoxy, 1-Boc-piperdin-4-ylmethoxy, piperdin-
		4-ylmethoxy, 1-methylpiperdin-4-yloxy,
		isopropoxy, methoxy and ethoxy;
	wherein R ²	is one or more substituents independently

wherein R^2 is one or more substituents independently selected from

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Η.
                halo,
                hvdroxy.
                amino.
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                C1_6-alkvl.
                C1-6-haloalkyl,
                C_{1-6}-alkoxy,
                C1_2-alkvlamino.
                aminosulfonyl,
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                C3-6-cycloalkyl,
                cvano.
                C1-2-hydroxyalkyl,
                nitro.
                Cola-alkenvl,
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                C_{2-3}-alkynyl,
                C1-6-haloalkoxy.
                C1-6-carboxvalkvl,
                5-6-membered heterocyclyl-C1-6-alkylamino,
                unsubstituted or substituted phenyl and
20
                unsubstituted or substituted 5-6 membered
                  heterocyclyl.
       preferably H, chloro, fluoro, bromo, amino, hydroxy,
          methyl, ethyl, propyl, oxo, dimethylamino,
          aminosulfonyl, cyclopropyl, cyano, hydroxymethyl,
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          nitro, propenyl, trifluoromethyl, methoxy, ethoxy,
          trifluoromethoxy, carboxymethyl,
          morpholinylethylamino, propynyl, unsubstituted or
          substituted phenyl and unsubstituted or substituted
          heteroaryl selected from thienyl, furanyl,
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                pyridyl, imidazolyl, and pyrazolyl,
           specifically chloro, fluoro, bromo, amino, hydroxy,
              methyl, ethyl, propyl, oxo, dimethylamino,
              aminosulfonyl, cyclopropyl, cyano, hydroxymethyl,
              nitro, propenyl, trifluoromethyl, methoxy, ethoxy,
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trifluoromethoxy, carboxymethyl,

morpholinylethylamino, propynyl, unsubstituted or substituted phenyl and unsubstituted or substituted heteroaryl selected from thienyl, furanyl,

pyridyl, imidazolyl, and pyrazolyl;

wherein R4 is selected from a direct bond, C1-4-alkyl, and

preferably a direct bond, ethyl, butyl, and $^{\prime\prime}$; wherein R^z is selected from C₁₋₂-alkyl, C₂₋₆-branched alkyl,

10 C_{2-4} -branched haloalkyl, amino- C_{1-4} -alkyl and C_{1-2} -alkylamino- C_{1-2} -alkyl,

wherein R^{\bullet} and R^{f} are independently selected from H and C_{1-2} haloalky1,

preferably trifluoromethyl: and

wherein R^7 is selected from H, C_{1-3} -alkyl, optionally substituted phenyl, optionally substituted phenyl- C_{1-3} alkyl, optionally substituted 4-6 membered

heterocyclyl, optionally substituted 4-6 membered heterocyclyl- C_1 - C_3 -alkyl, C_{1-3} -alkoxy- C_{1-2} -alkyl and C_{1-3} -alkoxy- C_{1-1} -alkoxy- C_{1-1} -alkoxy- C_{1-1} -alkyl;

provided R¹ is substituted with optionally substituted phenyloxy, optionally substituted 4-6 membered heterocyclyloxy, optionally substituted 4-6 membered heterocyclyl-C₁₋₄-alkoxy, optionally substituted 4-6 membered heterocyclylsulfonyl, optionally substituted

4-6 membered heterocyclylamino, optionally substituted
4-6 membered heterocyclylcarbonyl, optionally

30 substituted 4-6 membered heterocyclyl-C₁₋₄-

alkylcarbonyl, C_{1-3} -alkylamino- C_{1-3} -alkoxy, or C_{1-3} -alkylamino- C_{1-3} -alkoxy- C_{1-3} -alkoxy; further provided R is not 3-pyridyl when R^5 is CH_2 ;

and pharmaceutically acceptable isomers and derivatives thereof.

The invention also relates to compounds of Formula XII

XII

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wherein R¹ is selected from unsubstituted or substituted aryl, preferably phenyl, tetrahydronaphthyl, indanyl, indenyl, and naphthyl,

cycloalkyl, preferably cyclohexyl,

- 5-6 membered heteroaryl, preferably isoxazolyl, pyrazolyl, thiazolyl, thiadiazolyl, thienyl, pyridyl, pyrimidinyl, and pyridazinyl, and
- 9-10 membered bicyclic and 13-14 membered tricyclic heterocyclyl, preferably 1,2-dihydroquinolyl, 1,2,3,4-20 tetrahydro-isoquinolyl, isoquinolyl, quinolyl, indolyl, isoindolyl, 2,3-dihydro-1H-indolyl, naphthyridinyl, quinozalinyl, benzo[d]isothiazolyl, 2,3,4,4a,9,9a-hexahydro-1H-3-aza-fluorenyl, 5,6,7-trihydro-1,2,4-triazolo[3,4-a]isoquinolyl,
- 25 tetrahydroquinolinyl, indazolyl, 2,1,3-

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benzothiadiazolyl, benzodioxanyl, benzothienyl, benzofuryl, benzimidazolyl, benzoxazolyl and benzthiazolyl:

wherein substituted R1 is substituted with one or more substituents selected from halo, C1-6-alkvl, optionally substituted Case-cycloalkyl, optionally substituted phenyl, optionally substituted phenyl-C1-C4-alkylenyl, C1-2-haloalkoxy, optionally substituted 4-6 membered heterocyclyl-C1-C4-alkyl, optionally substituted 4-6 membered heterocyclyl-C2-C4-alkenyl, optionally substituted 4-6 membered heterocyclyl, optionally substituted phenyloxy, optionally substituted 4-6 membered heterocyclyloxy, optionally substituted 4-6 membered heterocyclyl-C1-C4-alkoxy, optionally substituted 4-6 membered heterocyclylsulfonyl, optionally substituted 4-6 membered heterocyclylamino, optionally substituted 4-6 membered heterocyclylcarbonyl, optionally substituted 5-6 membered heterocyclyl-C1-4-alkylcarbonyl, C1-2haloalkyl, C1-4-aminoalkyl, nitro, amino, hydroxy, oxo, cvano, aminosulfonyl, C1-2-alkylsulfonyl, halosulfonyl, C1-4-alkylcarbonyl, C1-3-alkylamino-C1-3-alkyl, C1-3alkylamino-C1-3-alkoxy, C1-3-alkylamino-C1-3-alkoxy-C1-3alkoxy, C1-4-alkoxycarbonyl, C1-4-alkoxycarbonylamino-C1-

4-alkyl, C₁₋₄-hydroxyalkyl, 0 and C₁₋₄-alkoxy, preferably bromo, chloro, fluoro, iodo, nitro, amino, cyano, aminoethyl, Boc-aminoethyl, hydroxy, oxo, aminosulfonyl, 4-methylpiperazinylsulfonyl, cyclohexyl, phenyl, phenylmethyl, morpholinylmethyl, 1-methylpiperazin-4-ylmethyl, 1-methylpiperazin-4-ylpiperazin-4-ylpropyl, morpholinylpropyl, piperidin-1-ylmethyl, 1-methylpiperidin-4-ylmethyl, 2-methyl-2-(1-methylpiperidin-4-vl)ethyl, morpholinylethyl, 1-(4-

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ethoxy;

morpholinv1)-2.2-dimethylpropyl, piperidin-4-ylethyl, 1-Boc-piperidin-4-vlethyl, piperidin-1-vlethyl, 1-Bocpiperidin-4-vlethyl, piperidin-4-vlmethyl, 1-Bocpiperidin-4-ylmethyl, piperidin-4-ylpropyl, 1-Bocpiperidin-4-vlpropyl, piperidin-1-vlpropyl, pyrrolidin-1-vlpropyl, pyrrolidin-2-vlpropyl, 1-Bocpyrrolidin-2-vlpropyl, pyrrolidin-1-vlmethyl, pyrrolidin-2-ylmethyl, 1-Boc-pyrrolidin-2-ylmethyl, pyrrolidinylpropenyl, pyrrolidinylbutenyl, fluorosulfonyl, methylsulfonyl, methylcarbonyl, Boc, piperidin-1-ylmethylcarbonyl, 4-methylpiperazin-1ylcarbonylethyl, methoxycarbonyl, aminomethylcarbonyl, dimethylaminomethylcarbonyl, 3-ethoxycarbonyl-2methyl-fur-5-yl, 4-methylpiperazin-1-yl, 4-methyl-1piperidyl, 1-Boc-4-piperidyl, piperidin-4-yl, 1methylpiperidin-4-vl, 1-methyl-(1,2,3,6tetrahydropyridyl), imidazolyl, morpholinyl, 4trifluoromethyl-1-piperidinyl, hydroxybutyl, methyl, ethyl, propyl, isopropyl, butyl, tert-butyl, secbutyl, trifluoromethyl, pentafluoroethyl, nonafluorobutyl, dimethylaminopropyl, 1,1di(trifluoromethyl)-1-hydroxymethyl, 1,1di(trifluoromethyl)-1-(piperidinylethoxy)methyl, 1,1di(trifluoromethyl)-1-(methoxyethoxyethoxy)methyl, 1hydroxyethyl, 2-hydroxyethyl, trifluoromethoxy, 1aminoethyl, 2-aminoethyl, 1-(N-isopropylamino)ethyl, 2-(N-isopropylamino) ethyl, dimethylaminoethoxy, 4chlorophenoxy, phenyloxy, azetidin-3-vlmethoxy, 1-Bocazetidin-3-ylmethoxy, pyrrol-2-ylmethoxy, 1-Bocpyrrol-2-ylmethoxy, pyrrol-1-ylmethoxy, 1-methylpvrrol-2-vlmethoxy, 1-isopropyl-pvrrol-2-vlmethoxy, 1-Boc-piperdin-4-ylmethoxy, piperdin-4-ylmethoxy, 1methylpiperdin-4-yloxy, isopropoxy, methoxy and

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wherein R2 is one or more substituents independently selected from н. halo. hvdroxv. amino, C1-6-alkyl, C1-6-haloalkyl, C1_6-alkoxy, C1-2-alkylamino, aminosulfonvl. C1-6-cycloalkyl, cvano. C₁₋₂-hydroxyalkyl, nitro, C2-1-alkenvl. C2-1-alkynyl, C1-6-haloalkoxy, C_{1-6} -carboxyalkyl, 5-6-membered heterocyclyl-C1-6-alkylamino, unsubstituted or substituted phenyl and

heterocyclyl,

preferably H, chloro, fluoro, bromo, amino, hydroxy,

methyl, ethyl, propyl, oxo, dimethylamino,
aminosulfonyl, cyclopropyl, cyano, hydroxymethyl,

nitro, propenyl, trifluoromethyl, methoxy, ethoxy, trifluoromethoxy, carboxymethyl, morpholinylethylamino, propynyl, unsubstituted or

unsubstituted or substituted 5-6 membered

30 substituted phenyl and unsubstituted or substituted heteroaryl selected from thienyl, fury, pyridyl,

imidazolyl, and pyrazolyl;

wherein R^e and R^f are independently selected from H and C_{1-2} haloalky1,

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preferably trifluoromethyl:

- wherein R^7 is selected from H, C_{1-3} -alkyl, optionally substituted phenyl, optionally substituted phenyl- C_{1-3} -alkyl, optionally substituted 4-6 membered heterocyclyl,
- 5 optionally substituted 4-6 membered heterocyclyl-C₁-C₃alkyl, C₁₋₃-alkoxy-C₁₋₂-alkyl and C₁₋₃-alkoxy-C₁₋₃-alkoxy-C₁₋₃-alkyl; and
 - wherein R^{20} is one or more substituents selected from halo, amino, hydroxy, C_{1-6} -alkyl, C_{1-6} -haloalkyl, C_{1-6} -alkoxy, optionally substituted heterocyclyl- C_{1-6} -alkoxy,
 - optionally substituted heterocyclyl-C₁₋₆-alkylamino, optionally substituted heterocyclyl-C₁₋₆-alkyl, C₁₋₆alkylamino-C₂₋₄-alkynyl, C₁₋₆-alkylamino-C₁₋₆-alkoxy, C₁₋₆alkylamino-C₁₋₆-alkoxy-C₁₋₆-alkoxy, and optionally
- 15 substituted heterocyclyl- C_{2-4} -alkynyl,
 - preferably chloro, fluoro, amino, hydroxy, methyl, ethyl,
 propyl, trifluoromethyl, dimethylaminopropynyl, 1 methylpiperdinylmethoxy, dimethylaminoethoxyethoxy,
 methoxy and ethoxy;
- 20 and pharmaceutically acceptable isomers and derivatives thereof.
 - A family of specific compounds of particular interest within Formula I consists of compounds and pharmaceutically-acceptable derivatives thereof as follows:
 - N-(4-Isopropylphenyl) (2-[(4-pyridylmethyl)amino](3-pyridyl)}carboxamide;
 - N-[3-(Isopropyl)phenyl](2-[(4-pyridylmethyl)amino](3-pyridyl)}carboxamide;
- 30 N-(3-Isoquinolyl)(2-[(4-pyridylmethyl)amino](3-pyridyl))carboxamide:
 - N-[4-Isopropylphenyl](2-[(2-(3-pyridyl)ethyl)amino](3-pyridyl)}carboxamide;

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N-[4-(tert-Buty1)phenyl](2-[(2-(3-pyridy1)ethy1)amino](3-
pyridy1))carboxamide;
N-[4-(Methylpropy1)phenyl](2-[(2-(3-pyridy1)ethy1)amino](3-
pyridy1))carboxamide;
(2-[(2-(3-Pyridy1)ethy1)amino](3-pyridy1))-N-[3-
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- (trifluoromethy1)phenyl]carboxamide;
 (2-[(4-Pyridylmethy1)amino](3-pyridyl)}-N-{4-[2,2,2-trifluoro-1-hydroxy-1-
 - (trifluoro-i-nydroxy-i(trifluoromethyl)ethyl]phenyl}carboxamide;
- 10 N-[5-(tert-Butyl)isoxazol-3-y1]{2-[(4pyridylmethyl)amino](3-pyridyl)}carboxamide;
 - $N-[5-(tert-Buty1)-1-methylpyrazol-3-y1]{2-[(4-$
 - pyridylmethyl)amino](3-pyridyl)}carboxamide; N-[4-(tert-Butyl)(1,3-thiazol-2-yl)](2-[(4-
- 15 pyridylmethyl)amino](3-pyridyl)}carboxamide;
 - N-[5-(tert-Butyl)(1,3,4-thiadiazol-2-yl)]{2-[(4
 - pyridylmethyl) amino] (3-pyridyl) } carboxamide; N-[4-(4-Hydroxybutyl) phenyl] (2-[(4-pyridylmethyl) amino] (3pyridyl) } carboxamide;
- 20 N-[2-(4-Chlorophenyl)ethyl]-2-[(pyridin-4-ylmethyl)amino](3-pyridyl)carboxamide;
 - 5-Bromo-N-[2-(4-chlorophenyl) ethyl]-2-[(pyridin-4-ylmethyl)amino](3-pyridyl)carboxamide;
 - N-[2-(4-Phenoxyphenyl)ethyl]-2-[(pyridin-4-
- 25 ylmethyl)amino](3-pyridyl)carboxamide; N-[2-(4-Methoxyphenyl)ethyl]-2-[(pyridin-4-
 - N-[2-(4-Methoxypheny1)ethy1]-2-[(pyridin-4ylmethyl)amino](3-pyridyl)carboxamide;
 - N-[2-(3,4-Dimethoxyphenyl)ethyl]-2-[(pyridin-4-ylmethyl)amino] (3-pyridyl)carboxamide;
- 30 N-[2-(4-Hydroxy-3-ethoxyphenyl)ethyl]-2-[(pyridin-4-ylmethyl)amino] (3-pyridyl)carboxamide;
 - N-[2-(4-Fluorophenyl)ethyl]-2-[(pyridin-4-ylmethyl)amino](3-pyridyl)carboxamide;

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- N-[2-(4-(tert-Butyl)phenyl)ethyl]-2-[(pyridin-4-ylmethyl)amino] (3-pyridyl)carboxamide;
- N-[2-(3-Fluorophenyl)ethyl]-2-[(pyridin-4-ylmethyl)amino]
 (3-pyridyl)carboxamide:
- 5 N-[2-(3-Chlorophenyl)ethyl]-2-[(pyridin-4-ylmethyl)amino] (3-pyridyl)carboxamide;
 - N-[2-(3-(Trifluoromethyl)phenyl)ethyl]-2-[(pyridin-4ylmethyl)amino] (3-pyridyl)carboxamide;
 - $\label{eq:n-interpolation} \verb|N-[2-(3-Ethoxyphenyl)] 2-[(pyridin-4-ylmethyl)] amino| \\$
- N-[2-(3,4-Dimethylphenyl)ethyl]-2-[(pyridin-4-

(3-pyridyl)carboxamide:

- ylmethyl)amino] (3-pyridyl)carboxamide; N-[2-(1,3-Benzodioxol-5-yl)ethyl]-2-[(pyridin-4ylmethyl)amino] (3-pyridyl)carboxamide;
- 15 N-[2-(4-Methylphenyl)ethyl]-2-[(pyridin-4-ylmethyl)amino]
 (3-pyridyl)carboxamide:
 - N-[2-(4-Hydroxyphenyl)ethyl]-2-[(pyridin-4-ylmethyl)amino]
 (3-pyridyl)carboxamide;
 - N-[2-(3,4-Dimethoxyphenyl)ethyl]-2-[(pyridin-4-ylmethyl)amino] (3-pyridyl)carboxamide;
 - N-[2-(4-Bromophenyl)ethyl]-2-[(pyridin-4-ylmethyl)amino] (3-pyridyl)carboxamide;
 - N-[2-(3,4-Dichlorophenyl)ethyl]-2-[(pyridin-4ylmethyl)amino] (3-pyridyl)carboxamide;
- 25 N-[2-(4-(Fluorosulfonyl)phenyl)ethyl]-2-[(pyridin-4ylmethyl)amino] (3-pyridyl)carboxamide;
 - N-[2-(3,5-(Dimethoxy)phenyl)ethyl]-2-[(pyridin-4ylmethyl)amino| (3-pyridyl)carboxamide:
- N-[2-(2,4-Dichlorophenyl)ethyl]-2-[(pyridin-430 ylmethyl)amino] (3-pyridyl)carboxamide;
 - N-[2-(2-Fluorophenyl)ethyl]-2-[(pyridin-4-ylmethyl)amino]
 (3-pyridyl)carboxamide;
 - N-[2-(2-Chloropheny1) ethy1]-2-[(pyridin-4-ylmethy1) amino]
 (3-pyridy1) carboxamide;

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- N-[2-(4-(Aminosulphonyl)phenyl)ethyl]-2-[(pyridin-4vlmethyl)aminol (3-pyridyl)carboxamide:
- N-[2-(2-Thienyl)ethyl]-2-[(pyridin-4-ylmethyl)amino] (3-pyridyl)carboxamide;
- 5 N-[2-(Pyridin-2-y1)ethyl]-2-[(pyridin-4-ylmethyl)amino] (3-pyridyl)}carboxamide;
 - N-[2-(Pyridin-3-y1)ethyl]-2-[(pyridin-4-ylmethyl)amino] (3-pyridyl)carboxamide;
- N-[2-(Pyridin-4-y1)ethy1]-2-[(pyridin-4-y1methy1)amino] (3pyridy1)carboxamide:
 - N-(4-Phenylbutyl)-2-[(pyridin-4-ylmethyl)amino] (3pyridyl)carboxamide;
 - N-(2-Hydroxy-3-phenoxypropyl)-2-[(pyridin-4-ylmethyl)amino]
 (3-pyridyl)carboxamide;
- 15 {6-Chloro-5-fluoro-2-[(4-pyridylmethyl)amino](3-pyridyl)}-N[4-(isopropyl)phenyl]carboxamide;
 - {5-Fluoro-2-[(4-pyridylmethyl)amino](3-pyridyl)}-N-[4-(isopropyl)phenyl]carboxamide;
 - 2-[(Pyridin-4-ylmethyl) amino]-N-[4-tert-butyl-3-(1,2,3,6tetrahydropyridin-4-vl) phenyl1(3-pyridyl) carboxamide;
 - N-(3,4-Dichlorophenyl)(6-[(2-morpholin-4-ylethyl)amino]-2[(4-pvridvlmethyl)amino](3-pvridvl)}carboxamide;
 - N-[4-(Morpholin-4-ylmethyl)phenyl]{2-[(4-pyridylmethyl)amino](3-pyridyl)}carboxamide;
- 25 N-(4-(2-[(tert-Butoxy)carbonylamino]ethyl)phenyl) (2-[(4-pyridylmethyl)amino](3-pyridyl))carboxamide;
 - N-[4-(2-Aminoethyl)phenyl]{2-[(4-pyridylmethyl)amino](3-pyridyl)}carboxamide;
 - N-[4-(tert-Butyl)-3-nitrophenyl]{2-[(2-
- 30 pyridylmethyl)amino](3-pyridyl)}carboxamide;
 - N-[3-Amino-4-(tert-buty1)pheny1]{2-[(2pyridylmethy1)amino](3-pyridy1)}carboxamide;
 - N-[4-(Isopropyl)phenyl]{2-[(2-pyridylmethyl)amino](3-pyridyl)}carboxamide;

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```
N-(3-Aminosulfonyl-4-chlorophenyl){2-[(4-
       pyridylmethyl)amino](3-pyridyl)}carboxamide;
    N-{3-[(4-Methylpiperazinyl)sulfonyl]phenyl}{2-[(4-
       pyridylmethyl)amino](3-pyridyl)}carboxamide;
    N-[4-(1,1,2,2,2-Pentafluoroethyl)phenyl]{2-[4-
       pyridylmethyl)amino](3-pyridyl)}carboxamide;
    N-[4-(1,1,2,2,3,3,4,4,4-Nonafluorobutyl) phenyl]{2-[(4-
       pyridylmethyl)aminol(3-pyridyl)}carboxamide;
    N-[4-(Isopropyl)phenyl]{2-[(2-(1,2,4-
10
       triazolyl)ethyl)amino](3-pyridyl)}carboxamide;
     (2-{[2-(2-Pvridvlamino)ethvl]amino}(3-pvridvl))-N-[3-
       (trifluoromethyl)phenyl]carboxamide;
     {2-[(1-(2-Pyridyl)pyrrolidin-3-yl)amino](3-pyridyl)}-N-[3-
       (trifluoromethyl)phenyl]carboxamide;
15
    2-[(Pvridin-4-vlmethvl)-amino]-N-(3-trifluoromethvl-phenvl)-
       nicotinamide
     {2-[(4-Pyridylmethyl)amino](3-pyridyl)}-N-(8-
       quinolvl)carboxamide hvdrochloride:
    N-[4-(4-Chlorophenoxy)phenyl] (2-[(4-pyridylmethyl)amino](3-
20
       pyridyl) } carboxamide hydrochloride:
     {2-[(4-Pyridylmethyl)amino](3-pyridyl)}-N-(2,3,4-
       trifluorophenyl)carboxamide hydrochloride;
    N-(2-Naphthyl){2-((4-pyridylmethyl)amino)(3-
       pyridyl)}carboxamide hydrochloride;
25
    N-(2-Phenoxyphenyl) {2-[(4-pyridylmethyl)amino](3-
       pyridyl) } carboxamide hydrochloride;
     {2-[(4-Pyridylmethyl)amino](3-pyridyl)}-N-(5,6,7,8-
       tetrahydronaphthyl) carboxamide hydrochloride;
    N-(2H-Benzo[3,4-d]1,3-dioxolen-5-yl){2-[(4-
30
       pyridylmethyl)amino](3-pyridyl)}carboxamide
       hydrochloride;
    N-Naphthyl (2-[(4-pyridylmethyl)amino](3-pyridyl))
```

carboxamide hydrochloride;

15

20

25

30

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```
N-[3-Benzylphenyl] {2-[(4-pyridylmethyl)amino](3-pyridyl)}
   carboxamide hydrochloride:
N-(Cyclohexylethyl) {2-[(4-pyridylmethyl)amino](3-pyridyl)}
   carboxamide hydrochloride:
N-(Cyclohexylethyl) {2-[(4-pyridylmethyl)amino](3-pyridyl)}
   carboxamide hydrochloride;
N-Indan-2-v1(2-[(4-pvridvlmethvl)amino](3-pvridvl)}
   carboxamide hydrochloride:
N-[4-(tert-Butyl)phenyl]{2-[(4-pyridylmethyl)amino](3-
   pyridyl) } carboxamide;
N-(4-sec-Butyl-phenyl)-2-[(pyridin-4-ylmethyl)-amino]-
   nicotinamide:
N-(4-Methylphenyl) {2-[(4-pyridylmethyl)amino](3-pyridyl)}
   carboxamide:
{2-[(4-Pyridylmethyl)amino](3-pyridyl)}-N-[4-
   trifluoromethoxy)phenyl] carboxamide;
N-(4-Ethylphenyl) {2-[(4-pyridylmethyl)amino](3-pyridyl)}
   carboxamide:
N-(4-Butylphenyl) (2-[(4-pyridylmethyl)amino](3-pyridyl)}
   carboxamide:
N-(4-Iodophenyl) {2-[(4-pyridylmethyl)aminol(3-pyridyl)}
   carboxamide:
N-[3-(Hydroxyethyl)phenyl]{2-[(4-pyridylmethyl)amino](3-
   pyridyl) } carboxamide;
N-(3-Ethylphenyl) {2-[(4-pyridylmethyl)amino](3-pyridyl)}
   carboxamide;
```

- Ethyl 2-methyl-5-[3-({2-[(4-pyridylmethyl)amino](3pyridyl)}carbonvlamino)phenyl|furan-3-carboxylate;
- N-(3-Phenylphenyl) {2-[(4-pyridylmethyl)amino](3-

pvridvl) {carboxamide;

- N-[4-Benzylphenyl] {2-[(4-pyridylmethyl)amino](3-pyridyl)}
 carboxamide:
 - N-(6-Ethyl(2-pyridyl))(2-[(4-pyridylmethyl)amino](3pyridyl)) carboxamide;

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```
N-(6-Propyl(2-pyridyl)){2-[(4-pyridylmethyl)amino](3-
        pyridyl)} carboxamide;
     N-[4-(tert-Butyl)(2-pyridyl)]{2-[(4-pyridylmethyl)amino](3-
        pvridvl) } carboxamide:
 5
    N-(3-Hydroxyphenyl) {2-[(4-pyridylmethyl)amino](3-pyridyl)}
        carboxamide:
    N-[4-(Methylethyl)(2-pyridyl)]{2-[(4-pyridylmethyl)amino](3-
        pyridyl)) carboxamide:
    N-[3,5-bis(Trifluoromethyl)phenyl]{2-[(4-
10
        pyridylmethyl)aminol(3-pyridyl)}carboxamide
        hydrochloride:
     N-[4-Chloro-3-(trifluoromethyl)phenyl]{2-[(4-
        pyridylmethyl)aminol(3-pyridyl)} carboxamide
        hvdrochloride:
15
    N-(3-Chlorophenyl) {2-[(2-(4-pyridyl)ethyl)amino](3-
        pyridyl) } carboxamide hydrochloride;
     N-(4-Phenoxyphenyl){2-[(2-(2-pyridyl)ethyl)amino](3-
        pyridyl)}carboxamide:
     2-[(Benzo[b]thiophen-3-vlmethyl)amino](3-pyridyl)}-N-(4-
20
        phenoxyphenyl) carboxamide;
     N-(4-Phenoxyphenv1)\{2-(2-(3-pyridy1)ethv1)amino\}(3-
       pyridyl) } carboxamide;
    N-[4-(Methylsulfonyl)phenyl]{2-[(4-pyridylmethyl)amino](3-
       pyridyl) } carboxamide:
25
    N-(1-Acetylindolin-6-yl){2-[(4-pyridylmethyl)amino](3-
        pyridyl) } carboxamide:
    N-Indolin-6-y1{2-[(4-pyridylmethyl)amino](3-
        pyridyl)}carboxamide;
    N-Indol-6-v1{2-[(4-pyridylmethyl)amino](3-
30
        pyridyl) } carboxamide;
    N-Indol-5-v1{2-[(4-pyridylmethyl)amino](3-
        pyridyl)}carboxamide;
    N-Indol-7-v1(2-[(4-pyridylmethyl)amino](3-
```

pyridyl) } carboxamide;

C3

100

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```
N-[3-(tert-Butyl)pyrazol-5-yl]{2-[(4-pyridylmethyl)aminol(3-
        pvridvl) {carboxamide:
     N-(3-Phenylpyrazol-5-yl){2-[(4-pyridylmethyl)amino](3-
        pyridyl) }carboxamide;
 5
     N-\{2-[2-(dimethylamino)ethoxy]-5-(tert-butyl)phenyl\}\{2-[(4-
        pyridylmethyl)aminol(3-pyridyl)}carboxamide;
     N-[4-(tert-Buty1)-3-(4-methylpiperaziny1)pheny1]{2-[(4-methylpiperaziny1)pheny1]}
        pyridylmethyl)aminol(3-pyridyl)}carboxamide;
     N-[3-(4-Methylpiperazinyl)phenyl]{2-[(4-
10
        pyridylmethyl)aminol(3-pyridyl)}carboxamide;
     N-[4-(4-Methylpiperazinyl)phenyl]{2-[(4-
        pyridylmethyl)aminol(3-pyridyl)}formamide:
     N-[1-(1-Methyl-(4-piperidyl))indolin-6-yl]{2-[(4-
        pyridylmethyl)amino](3-pyridyl)}carboxamide;
15
     N-[1-(1-Methyl-(4-piperidyl))] indolin-6-yl]{2-[(2-(3-
        pyridyl) ethyl) amino] (3-pyridyl) } carboxamide;
     N-[1-(2-Piperidvlethvl)indolin-6-vl]{2-[(4-
        pyridylmethyl)aminol(3-pyridyl)}carboxamide;
     N-[1-(2-Piperidylacetyl)indolin-6-yl]{2-[(4-
20
        pyridylmethyl)aminol(3-pyridyl)}carboxamide;
     N-[3.3-Dimethyl-1-(1-methyl(4-piperidyl))] indolin-6-yll(2-
        [(4-pyridylmethyl)amino](3-pyridyl)}carboxamide;
     N-(3,3-Dimethylindolin-6-yl){2-[(4-pyridylmethyl)amino](3-
        pyridyl) }carboxamide;
25
     N-[3-(1-Methyl-(4-piperidyl))indol-5-yl]{2-[(4-piperidyl)]}
        pyridylmethyl)aminol(3-pyridyl)}carboxamide:
     N-[4-(1,1-Dimethyl-3-morpholin-4-vlpropyl)phenyl]{2-[(4-(1,1-Dimethyl-3-morpholin-4-vlpropyl)phenyl]}
        pyridylmethyl)amino](3-pyridyl)}carboxamide;
     N-[4-(tert-Butyl)phenyl]{2-[({2-[(1-methyl(4-piperidyl))-
30
        methoxy](4-pyridyl)}methyl)amino](3-pyridyl)}carboxamide;
     N-(4-Bromo-2-fluorophenyl) {2-[(4-pyridylmethyl)amino](3-
        pyridyl) {carboxamide;
     N-[4-(tert-Butyl)phenyl](2-{[(2-chloro(4-
        pyridyl))methyl]amino}(3-pyridyl))carboxamide;
```

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```
{2-[({2-[3-(Dimethylamino)prop-1-ynyl](4-
        pyridyl) methyl) amino] (3-pyridyl) }-N-[4-(tert-
        butvl)phenvllcarboxamide:
     (2-{[(2-Methoxy(4-pyridyl))methyl]amino}(3-pyridyl))-N-[4-
 5
        (methylethyl)phenyllcarboxamide:
     N-{3-[3-(Dimethylamino)propyl]-5-(trifluoromethyl)phenyl}-
        {2-[(4-pyridylmethyl)amino](3-pyridyl)}carboxamide;
     N-[4-(tert-Butyl)-3-(3-piperid-1-ylpropyl)phenyl]{2-[(4-
        pyridylmethyl)aminol(3-pyridyl)}carboxamide;
10
     N-[4-(tert-Butvl)-3-(3-pyrrolidin-1-vlpropyl)phenyl]{2-[(4-
        pyridylmethyl)aminol(3-pyridyl)}carboxamide:
     N-[3-((1E)-4-Pvrrolidin-1-v]but-1-env]-4-(tert-
        butyl)phenyl]{2-[(4-pyridylmethyl)amino](3-
        pyridyl) } carboxamide;
15
     N-(4-(tert-Butyl)-3-(3-morpholin-4-vlpropyl)phenyl]{2-((4-
        pyridylmethyl)amino](3-pyridyl)}carboxamide;
     N-[1-(2-Morpholin-4-ylethyl) indol-6-yl]{2-[(4-ylethyl)]}
        pvridvlmethvl)aminol(3-pvridvl)}carboxamide:
     N-[4-(tert-Butyl)phenyl]{2-[(pyrimidin-4-ylmethyl)amino](3-
20
        pyridyl) } carboxamide:
     N-(4-Chlorophenyl) {2-[(pyrimidin-4-vlmethyl)amino](3-
        pyridyl) } carboxamide;
     {2-[(Pyrimidin-4-ylmethyl)amino](3-pyridyl)}-N-[3-
        (trifluoromethyl)phenyllcarboxamide;
25
     N-[4-(Isopropyl)phenyl]{4-[(4-pyridylmethyl)amino]pyrimidin-
        5-vl}carboxamide:
     (2-{[(2-{2-[2-(Dimethylamino)ethoxy]ethoxy}(4-
        pyridyl))methyl]amino}(3-pyridyl))-N-[4-(tert-
        butyl)phenyl]carboxamide;
30
     {2-[(4-Pvridvlmethyl)amino](3-pvridvl)}-N-{4-[2,2,2-
        trifluoro-1-(2-piperidylethoxy)-1-
```

(trifluoromethyl)ethyl)phenyl}carboxamide;

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```
(2-{[(2-{2-[2-(Dimethylamino)ethoxy]ethoxy}(4-
        pyridyl))methyl]amino}-6-fluoro(3-pyridyl))-N-[3-
        (trifluoromethyl)phenyllcarboxamide:
     N-[4-(tert-Butv1)phenv1]{6-fluoro-2-[(4-
 5
        pyridylmethyl)aminol(3-pyridyl)}carboxamide;
     (6-Fluoro-2-[(4-pyridylmethyl)aminol(3-pyridyl)}-N-[4-
        (isopropyl)phenyllcarboxamide:
     {6-Fluoro-2-[(4-pyridylmethyl)amino](3-pyridyl)}-N-[3-
        (trifluoromethyl)phenyl]carboxamide;
10
    N-(1-Bromo(3-isoguinolv1)) {6-fluoro-2-f(4-isoguinolv1)}
        pyridylmethyl)aminol(3-pyridyl)}-carboxamide:
    N-(4-Phenoxyphenyl) {2-[(4-pyridylmethyl)amino](3-
        pyridyl)}carboxamide hydrochloride:
    N-(4-Phenylphenyl) {2-[(4-pyridylmethyl)amino](3-
15
        pyridyl)}carboxamide hydrochloride;
    N-(3-Phenoxyphenyl) {2-[(4-pyridylmethyl)amino](3-
        pyridyl) } carboxamide hydrochloride;
    N-(4-Cyclohexylphenyl) {2-[(4-pyridylmethyl)amino](3-
        pyridyl) } carboxamide hydrochloride;
2.0
    N-(4-Imidazol-1-vlphenvl)(2-[(4-pvridvlmethvl)aminol(3-
        pyridyl) } carboxamide;
    N-(4-Morpholin-4-ylphenyl) {2-[(4-pyridylmethyl)amino](3-
        pyridyl) }carboxamide hydrochloride:
    N-(4-Cyanonaphthyl) {2-[(4-pyridylmethyl)amino](3-
25
        pyridyl) } carboxamide hydrochloride;
     {2-[(4-Pyridylmethyl)amino](3-pyridyl)}-N-[4-
        (trifluoromethyl)phenyl]carboxamide hydrochloride:
    Methyl-4-({2-[(4-pyridylmethyl)amino]-3-
        pyridyl}carbonylamino)benzoate hydrochloride;
30
    N-[4-(Isopropyl)phenyl]{2-[(4-quinolylmethyl)amino](3-
        pyridyl) { carboxamide:
```

N-[4-(tert-Butyl)phenyl]{2-[(6-quinolylmethyl)amino](3-

pyridyl) } carboxamide;

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```
(2-[(6-Quinolylmethyl)amino](3-pyridyl)}-N-[3-
(trifluoromethyl)phenyl]carboxamide;
N-(4-chlorophenyl)(3-[(4-pyridylmethyl)amino](2-
thienyl))carboxamide;
```

- 5 N-phenyl(3-[(4-pyridylmethyl)amino](2-thienyl))carboxamide;
 - N-(4-chlorophenyl)-3-[(4-pyridinylmethylene)amino]-4pyridinecarboxamide:
 - N-(4-chlorophenyl) {2-[(4-pyridylmethyl)amino](3pyridyl)}carboxamide;
- 10 N-(3,4-dichlorophenyl) (2-[(4-pyridylmethyl)amino](3pyridyl))-carboxamide;
 - N-(3-chlorophenyl) {2-[(4-pyridylmethyl)amino](3-pyridyl)}carboxamide;
- N-(4-chlorophenyl)(3-[(4-pyridylmethyl)amino](2-15 pyridyl))carboxamide:
 - N-(4-chlorophenyl) (3-[(6-quinolylmethyl)amino](2pvridvl))carboxamide:
 - N-(3,4-dichlorophenyl)(2-[(6-quinolylmethyl)amino](3-pyridyl)}-carboxamide;
- 20 N-(4-chlorophenyl)(6-methyl-2-[(4-pyridylmethyl)amino](3pyridyl))carboxamide;
 - N-(3,4-dichlorophenyl){6-methyl-2-[(4pyridylmethyl)amino](3-pyridyl)}carboxamide;
 - N-(3-fluoro-4-methylphenyl) {6-methyl-2-[(4-
- 25 pyridylmethyl)amino](3-pyridyl)}carboxamide;
 - N-(3,4-dichlorophenyl){6-chloro-2-[(4pyridylmethyl)amino](3-pyridyl)}carboxamide;
 - N-(4-chlorophenyl)(6-chloro-2-[(4-pyridylmethyl)amino](3-pyridyl))carboxamide;
- 30 (6-chloro-2-[(4-pyridylmethyl)amino](3-pyridyl)}-N-(3fluorophenyl)carboxamide;
 - N-(3-chlorophenyl)(6-chloro-2-[(4-pyridylmethyl)amino](3-pyridyl))carboxamide;

15

35

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```
N-(4-chlorophenyl) (3-[(4-pyridylmethyl)amino] (4-pyridyl) carboxamide:
```

- N-(3-fluoro-4-methylphenyl) {2-[(4-pyridylmethyl)amino](3-pyridyl)}carboxamide;
- 5 N-(4-chlorophenyl) {2-[(4-quinolylmethyl)amino](3pyridyl)}carboxamide:
 - N-(4-chlorophenyl) {2-[(5-quinolylmethyl)amino](3pyridyl)}carboxamide:
 - N-(4-chlorophenyl)(2-[(4-pyridylethyl)amino]-5-(3-thienyl)-(3-pyridyl))carboxamide;
- N-(4-chlorophenyl) {5-(4-methoxyphenyl)-2-[(4
 - pyridylmethyl)amino]-(3-pyridyl))carboxamide; and
 - N-(4-chlorophenyl)(5-bromo-2-[(4-pyridylmethyl)amino]-(3-pyridyl))carboxamide.
 - A family of specific compounds of particular interest within Formula II' consists of compounds and pharmaceutically-acceptable derivatives thereof as follows:
- - N-(4-tert-Butyl-phenyl)-2-{[2-(1-isopropyl-azetidin-3ylmethoxy)-pyridin-4-ylmethyl]-amino)-nicotinamide;
- 25 2-[(2,3-Dihydro-benzofuran-5-ylmethyl)-amino]-N-(4-[1methyl-1-(1-methyl-piperidin-4-yl)-ethyl]-phenyl)nicotinamide;
 - N-(1-Acetyl-3,3-dimethyl-2,3-dihydro-1H-indol-6-yl)-2-[(2,3-dihydro-benzofuran-5-ylmethyl)-amino]-nicotinamide;
- - 2-[(2,3-Dihydro-benzofuran-5-ylmethyl)-amino]-N-[3,3 dimethyl-1-(1-methylpiperidin-4-ylmethyl)-2,3-dihydro1H-indol-6-yl]-nicotinamide;

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nicotinamide:

```
N-(1-Acetvl-3.3-dimethvl-2.3-dihvdro-1H-indol-6-vl)-2-({2-
          [2-(1-methyl-piperidin-4-yl)-ethoxyl-pyridin-4-
          vlmethvl}-amino)-nicotinamide:
     2-({2-[2-(1-Methyl-piperidin-4-yl)-ethoxy]-pyridin-4-
 5
          vlmethvl}-amino)-N-(3-trifluoromethvl-phenvl)-
          nicotinamide:
     N-(4-tert-Butyl-phenyl)-2-{[2-ethylpyridin-4-ylmethyl]-
          amino}-nicotinamide:
     N-(4-tert-Butvl-phenyl)-2-({2-[2-(1-methyl-pyrrolidin-2-yl)-
10
          ethoxyl-pyridin-4-vlmethyl}-amino)-nicotinamide;
     2-({2-[2-(1-Methyl-pyrrolidin-2-yl)-ethoxy]-pyridin-4-
          vlmethyl}-amino)-N-(4-pentafluoroethyl-phenyl)-
          nicotinamide:
     N-(4-Pentafluoroethyl-phenyl)-2-{[2-(2-pyrrolidin-1-yl-
15
          ethoxy)-pyridin-4-ylmethyl]-amino}-nicotinamide;
     N-(4-\text{tert-Butyl-phenyl})-2-([2-(2-\text{pyrrolidin-1-vl-ethoxy})-
          pyridin-4-vlmethyl]-amino}-nicotinamide;
     N-[3-(4-Boc-piperazin-1-ylmethyl)-5-trifluoromethyl-phenyl]-
          2-[(pyridin-4-ylmethyl)-amino]-nicotinamide;
20
    N-[3-(4-Boc-piperazine-1-carbonvl)-5-trifluoromethyl-
          phenyl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide;
    N-[3-(4-Boc-piperazine-1-carbonyl)-5-trifluoromethyl-
          phenv11-2-(2-pvridin-4-vl-ethvlamino)-nicotinamide:
    N-[3-(4-Methyl-piperazin-1-ylmethyl)-4-pentafluoroethyl-
25
          phenyl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide;
     N-[3-(4-Boc-piperazin-1-vlmethyl)-4-pentafluoroethyl-
          phenyl]-2-[(pyridin-4-vlmethyl)-amino]-nicotinamide;
     2-{[2-(1-Methyl-piperidin-4-ylmethoxy)-pyridin-4-ylmethyl]-
          amino}-N-(4-trifluoromethyl-phenyl)-nicotinamide;
30
    N-(4-tert-Butvl-phenvl)-2-{[2-(1-methvl-piperidin-4-
          ylmethoxy)-pyridin-4-ylmethyl]-amino}-nicotinamide;
     2-({2-[3-(1-Methyl-piperidin-4-yl)-propoxy]-pyridin-4-
          vlmethyl}-amino)-N-(4-pentafluoroethyl-phenyl)-
```

10

25

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aminol-nicotinamide:

- N-(1-Acetyl-3,3-dimethyl-2,3-dihydro-1H-indol-6-yl)-2-[(2-methoxy-pyridin-4-vlmethyl)-aminol-nicotinamide;
- N-[3,3-Dimethyl-1-(1-methyl-piperidin-4-yl)-2,3-dihydro-1Hindol-6-yl]-2-[(2-methoxy-pyridin-4-ylmethyl)-amino]nicorinamide:
- N-(1-Boc-3,3-dimethyl-2,3-dihydro-1H-indol-6-yl)-2-[(2-methoxy-pyridin-4-ylmethyl)-aminol-nicotinamide:
- N-[3,3-Dimethyl-1-(1-Boc-piperidin-4-ylmethyl)-2,3-dihydro-1H-indol-6-yl]-2-[(2-methoxy-pyridin-4-ylmethyl)-
- N-[3,3-Dimethyl-1-(1-methyl-piperidin-4-yl)-2,3-dihydro-1Hindol-6-yl]-2-[(2-methoxy-pyridin-4-ylmethyl)-amino]nicotinamide:
- N-[1-(2-Dimethylamino-acetyl)-3,3-dimethyl-2,3-dihydro-1Hindol-6-yl]-2-[(2-methoxy-pyridin-4-ylmethyl)-amino]nicotinamide:
 - N-[1-(2-Dimethylamino-acetyl)-3,3-dimethyl-2,3-dihydro-1H-indol-6-yl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide;
- 20 2-[(2-Methoxy-pyridin-4-ylmethyl)-amino]-N-[3-(1-Boc-piperidin-4-ylmethoxy)-5-trifluoromethyl-phenyl]-nicotinamide;
 - N-[3,3-Dimethyl-1-(1-Boc-pyrrolidin-2-ylmethoxy)-2,3dihydro-1H-indol-6-yl]-2-[(2-methoxy-pyridin-4ylmethyl)-amino]-nicotinamide;
 - N-[3,3-Dimethyl-1-(2-Boc-amino-acetyl)-2,3-dihydro-1H-indol-6-yl]-2-[(2-methoxy-pyridin-4-ylmethyl)-amino]nicotinamide;
- N-[3,3-Dimethyl-1-(2-Boc-amino-acetyl)-2,3-dihydro-1H-indol-30 6-yl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide;
 - 2-[(2-Methoxy-pyridin-4-ylmethyl)-amino]-N-[3-(1-methylpyrrolidin-2-ylmethoxy)-5-trifluoromethyl-phenyl]nicotinamide:

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nicotinamide:

nicotinamide:

- 2-[(2-Methoxy-pyridin-4-vlmethyl)-amino]-N-[3-(1-Bocpiperidin-4-vlmethvl)-5-trifluoromethvl-phenvl]nicotinamide:
- 2-[(2-Methoxy-pyridin-4-ylmethyl)-amino]-N-[3-(4-Bocpiperazin-1-vlmethvl)-5-trifluoromethvl-phenvl]nicotinamide:
- 2-{[2-(3-Morpholin-4-yl-propoxy)-pyridin-4-ylmethyl]-amino}-N-(4-pentafluoroethyl-phenyl)-nicotinamide:
- (S) 2-{[2-(1-Methyl-pyrrolidin-2-ylmethoxy)-pyridin-4vlmethyll-amino}-N-(4-pentafluoroethyl-phenyl)-
- $N-(3-tert-Butvl-isoxazol-5-vl)-2-\{[2-(3-morpholin-4-vl$ propoxy)-pyridin-4-ylmethyl]-amino}-nicotinamide;
- N-(1-Acetyl-3,3-dimethyl-2,3-dihydro-1H-indol-6-yl)-2-{[2-15 (3-morpholin-4-yl-propylamino)-pyridin-4-ylmethyl]amino}-nicotinamide:
 - N-(4-tert-Butyl-phenyl)-2-{[2-(3-morpholin-4-yl-propoxy)pyridin-4-ylmethyl]-amino}-nicotinamide;
 - N-(4-tert-Butyl-phenyl)-2-([2-(2-morpholin-4-yl-ethoxy)pyridin-4-vlmethyll-amino}-nicotinamide;
 - 2-{[2-(2-Morpholin-4-yl-ethoxy)-pyridin-4-ylmethyl]-amino}-N-(4-trifluoromethyl-phenyl)-nicotinamide;
 - 2-{[2-(2-Morpholin-4-vl-ethoxy)-pvridin-4-vlmethvl]-amino}-N-(3-trifluoromethyl-phenyl)-nicotinamide;
- 25 2-{[2-(2-Morpholin-4-yl-ethoxy)-pyridin-4-ylmethyl]-amino}-N-(4-pentafluoroethyl-phenyl)-nicotinamide;
 - $N-(3-tert-Butyl-isoxazol-5-yl)-2-\{[2-(2-morpholin-4-yl$ ethoxy)-pyridin-4-ylmethyl]-amino}-nicotinamide;
- N-(1-Acetvl-3,3-dimethvl-2,3-dihvdro-1H-indol-6-vl)-2-{[2-30 (2-morpholin-4-vl-ethoxy)-pyridin-4-vlmethyl]-amino}-
 - $N-(1-Acetyl-3,3-dimethyl-2,3-dihydro-1H-indol-6-yl)-2-{[2-$ (1-methyl-piperidin-4-yloxy)-pyridin-4-ylmethyl]amino}-nicotinamide:

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nicotinamide:

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2-{[2-(1-Methyl-piperidin-4-vloxy)-pyridin-4-vlmethyl]-
          amino}-N-(4-trifluoromethyl-phenyl)-nicotinamide;
     2-{[2-(1-Methyl-piperidin-4-vloxy)-pyridin-4-vlmethyll-
          amino)-N-(4-pentafluoroethyl-phenyl)-nicotinamide:
    2-{[2-(1-Methyl-piperidin-4-vloxy)-pyridin-4-vlmethyl]-
          amino}-N-(4-tert-butyl-phenyl)-nicotinamide;
     (R) N-(4-tert-Butyl-phenyl)-2-{[2-(1-methyl-pyrrolidin-2-
          vlmethoxy) -pyridin-4-vlmethyll-amino}-nicotinamide;
     (R) N-[3-(1-Boc-pyrrolidin-2-vlmethoxy)-5-trifluoromethyl-
10
          phenyl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide;
     (R) N-[3-(1-Methyl-pyrrolidin-2-vlmethoxy)-5-
          trifluoromethyl-phenyl]-2-[(pyridin-4-ylmethyl)-
          aminol-nicotinamide:
    N-[3-(1-Methyl-piperidin-4-yloxy)-5-trifluoromethyl-phenyl]-
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          2-[(pvridin-4-vlmethvl)-amino]-nicotinamide;
    N-[3-(1-Methyl-piperidin-4-ylmethyl)-5-trifluoromethyl-
          phenyl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide;
    N-[3-tert-Butvl-4-(1-Boc-pvrrolidin-2-vlmethoxy)-phenvl]-2-
          [(pyridin-4-vlmethyl)-amino]-nicotinamide;
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    N-(3,3-Dimethyl-2,3-dihydro-benzofuran-6-yl)-2-{[2-(1-
          methyl-piperidin-4-vlmethoxy)-pyridin-4-vlmethyll-
          amino}-nicotinamide;
     2-({2-[3-(1-Methyl-piperidin-4-yl)-propoxy]-pyridin-4-
          ylmethyl}-amino)-N-(4-trifluoromethyl-phenyl)-
2.5
          nicotinamide:
     2-({2-[3-(1-Methyl-piperidin-4-yl)-propoxy]-pyridin-4-
          vlmethvl}-amino)-N-(3-trifluoromethvl-phenvl)-
          nicotinamide;
     2-({2-[3-(1-Methyl-piperidin-4-yl)-propoxy]-pyridin-4-
30
          ylmethyl}-amino)-N-(4-tert-butyl-phenyl)-nicotinamide;
     2-({2-[3-(1-Methyl-piperidin-4-yl)-propoxy]-pyridin-4-
          ylmethyl}-amino)-N-(3-tert-butyl-isoxazol-5-yl)-
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- N-(3,3-Dimethyl-2,3-dihydro-1H-indol-6-yl)-2-({2-[3-(1-methyl-piperidin-4-yl)-propoxy]-pyridin-4-ylmethyl)-amino)-nicotinamide:
- 2-[(Pyridin-4-ylmethyl)-amino]-N-(3,9,9-trimethyl-
- 5 2,3,4,4a,9,9a-hexahydro-1H-3-aza-fluoren-6-yl)nicotinamide;
 - N-[3,3-Dimethyl-1-(1-Boc-piperidin-4-ylmethyl)-2,3-dihydro-1H-indol-6-yl]-2-[(pyridin-4-ylmethyl)-amino]nicotinamide:
- 10 N-(4-Imidazol-1-ylphenyl) {2-[(4-pyridylmethyl)amino](3-pyridyl))carboxamide hydrochloride;
 - N-[3,3-Dimethyl-1-(1-methyl-piperidin-4-ylmethyl)-2,3dihydro-1H-indol-6-yl]-2-[(pyridin-4-ylmethyl)-amino]nicotinamide;
- 15 2-([2-(1-Methyl-piperidin-4-ylmethoxy)-pyridin-4-ylmethyl]amino)-N-(4-pentafluoroethyl-phenyl)-nicotinamide;
 - N-(3-tert-Butyl-isoxazol-5-yl)-2-{[2-(1-methyl-piperidin-4-ylmethoxy)-pyridin-4-ylmethyl]-amino}-nicotinamide;
- N-(1-Acetyl-3,3-dimethyl-2,3-dihydro-1H-indol-6-yl)-2-([2-20 (1-methyl-piperidin-4-ylmethoxy)-pyridin-4-ylmethyl]amino)-nicotinamide;
 - N-(4-tert-Butyl-phenyl)-2-([2-(3-morpholin-4-ylpropylamino)-pyrimidin-4-ylmethyl]-amino)nicotinamide;
- 25 2-{[2-(3-Morpholin-4-yl-propylamino)-pyrimidin-4-ylmethyl]amino)-N-(4-pentafluoroethyl-phenyl)-nicotinamide;
 - 2-{[2-(3-Morpholin-4-y1-propylamino)-pyrimidin-4-y1methy1]amino)-N-(3-trifluoromethyl-phenyl)-nicotinamide;
 - $N-(4-\texttt{tert-Butyl-phenyl})-2-(\{2-[2-(1-\texttt{methyl-pyrrolidin-}2-\texttt{yl})-1-(1-\texttt{methyl-p$
- 30 ethylamino]-pyrimidin-4-ylmethyl]-amino)-nicotinamide;
 N-(1-Acetyl-3,3-dimethyl-2,3-dihydro-1H-indol-6-yl)-2-((2[2-(1-methyl-pyrrolidin-2-yl)-ethylamino]-pyrimidin-4ylmethyl)-amino)-nicotinamide;

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N-(4-Phenoxyphenyl)(2-[(4-pyridylmethyl)aminol(3-
     pyridyl) carboxamide hydrochloride:
2-{[2-(1-Methyl-piperidin-4-vlmethoxy)-pyridin-4-vlmethyl]-
     amino -N-[3-(1-methyl-piperidin-4-yl)-5-
     trifluoromethyl-phenyll-nicotinamide:
N-(3-tert-Butyl-isoxazol-5-yl)-2-{[2-(1-methyl-piperidin-4-
     ylmethoxy)-pyridin-4-ylmethyl]-amino}-nicotinamide;
N-[3-(1-Boc-azetidin-3-vlmethoxy)-5-trifluoromethyl-phenyl]-
     2-[(pvridin-4-vlmethvl)-aminol-nicotinamide:
2-[(2-Methoxy-pyridin-4-vlmethyl)-amino]-N-[3-(1-Boc-
     azetidin-3-ylmethoxy)-5-trifluoromethyl-phenyl]-
     nicotinamide:
N-(3,3-Dimethylindolin-6-vl)(2-[(4-pyridylmethyl)aminol(3-
  pyridyl)}carboxamide phosphate salt;
N-(4-Morpholin-4-ylphenyl){2-[(4-
  pyridylmethyl)amino](3-pyridyl)}carboxamide
  hvdrochloride:
N-(4-Cyanonaphthyl) {2-[(4-pyridylmethyl)amino](3-
  pvridvl) } carboxamide, hvdrochloride:
{2-[(4-Pyridylmethyl)amino](3-pyridyl)}-N-[4-
   (trifluoromethyl)phenyl]carboxamide hydrochloride:
Methyl-({2-[(4-pyridylmethyl)amino]-3-
     pyridyl}carbonylamino)benzoate, hydrochloride;
2-[(Pyridin-4-vlmethyl)-amino]-N-(2,2,4-trimethyl-3,4-
     dihydro-2H-benzo[1,4]oxazin-6-yl)-nicotinamide;
N-(4-Acetyl-2,2-dimethyl-3,4-dihydro-2H-benzo[1,4]oxazin-6-
     yl)-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide;
N-(2,2-Dimethyl-3-oxo-3,4-dihydro-2H-benzo[1,4]oxazin-6-yl)-
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2-[(pvridin-4-vlmethvl)-amino]-nicotinamide:

2-{[2-(1-Benzhydry1-azetidin-3-yloxy)-pyridin-4-ylmethy1]amino)-N-(4-tert-buty1-pheny1)-nicotinamide;
N-(4,4-Dimethy1-1-oxo-1,2,3,4-tetrahydro-isoquinolin-7-y1)2-[(pyridin-4-ylmethy1)-amino)-nicotinamide;

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- N-(4-tert-Butyl-phenyl)-2-({2-[2-(1-methyl-piperidin-4-yl)-ethoxyl-pyridin-4-ylmethyl}-amino)-nicotinamide:
- N-(3-tert-Butyl-isoxazol-5-yl)-2-({2-[2-(1-methyl-piperidin-4-yl)-ethoxy}-pyridin-4-ylmethyl}-amino)-nicotinamide;
- 5 N-(3-trifluoromethylphenyl)-2-({2-[2-(1-methyl-piperidin-4-yl)-ethoxy]-pyridin-4-ylmethyl}-amino)-nicotinamide;
 - 2-[(2,3-Dihydro-benzofuran-6-ylmethyl)-amino]-N-[3-(1-Bocpyrrolidin-2-ylmethoxy)-4-pentafluoroethyl-phenyl]nicotinamide;
- 10 N-[3-(1-Methyl-piperidin-4-ylmethoxy)-4-pentafluoroethyl-phenyl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide hydrochloride;
 - (R) N-[3-(2-Hydroxy-3-pyrrolidin-1-yl-propoxy)-4pentafluoroethyl-phenyl]-2-[(pyridin-4-ylmethyl)amino]-nicotinamide;
 - (S) N-[3-(2-Hydroxy-3-pyrrolidin-1-yl-propoxy)-4pentafluoroethyl-phenyl]-2-[(pyridin-4-ylmethyl)amino]-nicotinamide;
 - N-[4-tert-Butyl-3-(1-methyl-piperidin-4-ylmethoxy)-phenyl]2-[(pyridin-4-ylmethyl)-amino]-nicotinamide;
 - N-[3-(1-Methyl-piperidin-4-ylmethoxy)-4-pentafluoroethylphenyl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide;
 - N-[4-Pentafluoroethyl-3-(2-piperidin-1-yl-ethoxy)-phenyl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide;
- - N-(4-Imidazol-1-ylphenyl) (2-[(4-pyridylmethyl) amino] (3-pyridyl)) carboxamide hydrochloride;
- N-(3,3-Dimethyl-2,3-dihydro-benzofuran-6-yl)-2-[(pyridin-4-30 ylmethyl)-amino]-nicotinamide hydrochloride;
 - 2-[(Pyridin-4-ylmethyl)-amino]-N-(4-tert-butyl-phenyl)nicotinamide hydrochloride;
 - N-[4-Trifluoromethyl-3-(2-piperidin-1-yl-ethoxy)-phenyl]-2[(pyridin-4-ylmethyl)-amino]-nicotinamide;

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- (S) N-[3-(1-Boc-pyrrolidin-2-ylmethoxy)-4-pentafluoroethylphenyl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide;
- (R) N-[3-(1-Boc-pyrrolidin-2-ylmethoxy)-4-trifluoromethylphenyl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide;
- 5 (R) N-[3-(1-Boc-pyrrolidin-2-ylmethoxy)-4-pentafluoroethyl-phenyl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide;
 - N-(4-tert-Butyl-phenyl)-2-([2-(1-methyl-piperidin-4-yloxy)-pyridin-4-ylmethyl-amino)-nicotinamide:
 - N-(3-Trifluoromethyl-phenyl)-2-{[2-(1-methyl-piperidin-4yloxy)-pyridin-4-vlmethyl]-amino}-nicotinamide;
 - N-(3-tert-Butyl-isoxazol-5-yl)-2-([2-(1-methyl-piperidin-4vloxy)-pyridin-4-vlmethyl]-amino)-nicotinamide:
 - N-[3-(3-Piperidin-1-yl-propyl)-5-trifluoromethyl-phenyl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide;
- 15 N-[3-(3-Morpholin-4-yl-propyl)-5-trifluoromethyl-phenyl]-2[(pvridin-4-vlmethyl)-amino]-nicotinamide;
 - 2-[(2-Methoxy-pyridin-4-ylmethyl)-amino]-N-[3-(1-Bocpiperidin-4-yloxy)-5-trifluoromethyl-phenyl]nicotinamide:
- 20 N-(3,3-Dimethylindolin-6-yl) {2-[(4-pyridylmethyl)amino](3-pyridyl)}carboxamide edisylate;
 - N-{4-tert-Butyl-3-[2-(1-Boc-piperidin-4-yl)-ethyl]-phenyl}2-[(pyridin-4-ylmethyl)-amino]-nicotinamide;
 - N-[4-tert-Butyl-3-(1-methyl-azetidin-3-ylmethoxy)-phenyl]-2[(pvridin-4-ylmethyl)-amino]-nicotinamide:
 - N-(3,3-Dimethyl-1,1-dioxo-2,3-dihydro-1H-1λbenzo[d]isothiazol-6-yl)-2-[(pyridin-4-ylmethyl)amino]-nicotinamide:
 - N-[1,1,4,4-Tetramethyl-1,2,3,4-tetrahydro-naphth-6-yl]-2-
- 30 [(pyridin-4-ylmethyl)-amino]-nicotinamide; N-{4-[1-Methyl-1-(1-methyl-piperidin-4-yl)-ethyl]-phenyl}-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide;
 - 2-[(2-Methoxy-pyridin-4-ylmethyl)-amino]-N-{4-[1-methyl-1-(1-methyl-piperidin-4-yl)-ethyl]-phenyl}-nicotinamide;

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- N-(3,3-Dimethyl-2,3-dihydro-benzofuran-6-yl)-2-[(pyridin-4vlmethyl)-aminol-nicotinamide;
- N-(3,3-Dimethyl-2,3-dihydro-1H-indol-6-yl)-2-({2-[2-(1-methyl-piperidin-4-yl)-ethoxy]-pyridin-4-ylmethyl}aminol-nicotinamide:
- 2-[(Pyridin-4-ylmethyl)-amino]-N-[3-(2-pyrrolidin-1-yl-ethoxy)-4-trifluoromethyl-phenyl]-nicotinamide hydrochloride:
- N-(2,2-Dimethyl-3,4-dihydro-2H-benzo[1,4]oxazin-6-yl)-210 [(pyridin-4-ylmethyl)-amino]-nicotinamide;
 - N-(4,4-Dimethyl-1,2,3,4-tetrahydro-isoquinolin-7-yl)-2[(pvridin-4-vlmethyl)-aminol-nicotinamide;
 - N-(3,3-Dimethyl-2,3-dihydro-1H-indol-6-yl)-2-([2-(1-methyl-piperidin-4-ylmethoxy)-pyridin-4-ylmethyl]-amino)-nicotinamide:
 - N-(3,3-Dimethylindolin-6-yl) (2-[(4-pyridylmethyl)amino](3pyridyl))carboxamide hydrochloride;
 - N-(3,3-Dimethyl-1-piperidin-4-yl-2,3-dihydro-1H-indol-6-yl)-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide;
- 20 N-(3,3-dimethyl-2,3-dihydro-1H-indol-6-yl)-2-({2-[2-(1-methyl-pyrrolidin-2-yl)-ethylamino]-pyrimidin-4-ylmethyl}-amino)-nicotinamide;
 - N-(3,3-dimethyl-2,3-dihydro-1H-indol-6-yl)-2-[(2-methoxy-pyridin-4-ylmethyl)-amino]-nicotinamide;
- N-[3,3-Dimethyl-1-(piperidin-4-ylmethyl)-2,3-dihydro-1Hindol-6-yl]-2-[(2-methoxy-pyridin-4-ylmethyl)-amino]nicotinamide:
 - N-(3,3-Dimethyl-1-piperidin-4-yl-2,3-dihydro-1H-indol-6-yl)2-[(2-methoxy-pyridin-4-ylmethyl)-amino]-nicotinamide;
- 30 2-[(2-Methoxy-pyridin-4-ylmethyl)-amino]-N-[3-(piperidin-4-ylmethoxy)-5-trifluoromethyl-phenyl]-nicotinamide;
 - N-[3,3-Dimethyl-1-(pyrrolidin-2-ylmethoxy)-2,3-dihydro-1Hindol-6-yl]-2-[(2-methoxy-pyridin-4-ylmethyl)-amino]nicotinamide:

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- 2-[(2-Methoxy-pyridin-4-ylmethyl)-amino]-N-[3-(piperazin-1ylmethyl)-5-trifluoromethyl-phenyl]-nicotinamide;
- N-(3,3-dimethyl-2,3-dihydro-1H-indol-6-yl)-2-({2-(2-morpholin-4-yl-ethoxy)-pyridin-4-ylmethyl}-amino}-nicotinamide:
- N-(3,3-dimethyl-2,3-dihydro-1H-indol-6-yl)-2-({2-(2-morpholin-4-yl-propylamino)-pyridin-4-ylmethyl}amino)-nicotinamide hydrochloride;
- N-(3,3-dimethyl-2,3-dihydro-1H-indol-6-yl)-2-{[2-(1-methyl-10 piperidin-4-yloxy)-pyridin-4-ylmethyl]-amino}nicotinamide:
 - N-(3,3-dimethyl-2,3-dihydro-1H-indol-6-yl)-2-{{2-(2morpholin-4-yl-propoxy)-pyridin-4-ylmethyl}-amino}nicotinamide;
- 15 N-(4-Pentafluoroethyl-phenyl)-2-((pyrimidin-4-ylmethyl)aminol-nicotinamide:
 - 2-{[2-(Azetidin-3-yloxy)-pyridin-4-ylmethyl]-amino}-N-(4tert-butyl-phenyl)nicotinamide;
 - N-(2,3,3-Trimethyl-1,1-dioxo-2,3-dihydro-1H-1\(\lambda\)
 benzo[d]isothiazol-6-yl)-2-[(pyridin-4-ylmethyl)aminol-benzamide:
 - N- (4,4-Dimethyl-1-oxo-1,2,3,4-tetrahydro-isoquinolin-7-yl) -2-[(pyridin-4-ylmethyl)-amino]-nicotinamide hydrochloride:
- 25 N-[3,3-Dimethyl-1,1-dioxo-2-(2-piperidin-1-yl-ethyl)-2,3-dihydro-1H-1λ'-benzo[d]isothiazol-6-yl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide; and
 - N-[2-(2-Dimethylamino-ethyl)-3,3-dimethyl-1,1-dioxo-2,3-dihydro-1H-1\lambda'-benzo[d]isothiazol-6-yl]-2-[(pyridin-4-ylmethyl)-aminol-nicotinamide.

Indications

Compounds of the present invention would be useful for, but not limited to, the prevention or treatment of angiogenesis related diseases. The compounds of the

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invention have kinase inhibitory activity, such as VEGFR/KDR inhibitory activity. The compounds of the invention are useful in therapy as antineoplasia agents or to minimize deleterious effects of VEGF.

Compounds of the invention would be useful for the treatment of neoplasia including cancer and metastasis, including, but not limited to: carcinoma such as cancer of the bladder, breast, colon, kidney, liver, lung (including small cell lung cancer), esophagus, gall-bladder, ovary,

pancreas, stomach, cervix, thyroid, prostate, and skin (including squamous cell carcinoma); hematopoietic tumors of lymphoid lineage (including leukemia, acute lymphocitic leukemia, acute lymphoblastic leukemia, B-cell lymphoma, Tcell-lymphoma, Hodgkin's lymphoma, non-Hodgkin's lymphoma,

hairy cell lymphoma and Burkett's lymphoma); hematopoietic

tumors of myeloid lineage (including acute and chronic myelogenous leukemias, myelodysplastic syndrome and promyelocytic leukemia); tumors of mesenchymal origin (including fibrosarcoma and rhabdomyosarcoma, and other sarcomas, e.g. soft tissue and bone); tumors of the central and peripheral nervous system (including astrocytoma. neuroblastoma, glioma and schwannomas); and other tumors (including melanoma, seminoma, teratocarcinoma, osteosarcoma, xenoderoma pigmentosum, keratoctanthoma,

Preferably, the compounds are useful for the treatment of neoplasia selected from lung cancer, colon cancer and breast cancer.

thyroid follicular cancer and Kaposi's sarcoma).

The compounds also would be useful for treatment of 30 ophthalmological conditions such as corneal graft rejection. ocular neovascularization, retinal neovascularization including neovascularization following injury or infection, diabetic retinopathy, retrolental fibroplasia and neovascular glaucoma; retinal ischemia; vitreous hemorrhage;

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ulcerative diseases such as gastric ulcer; pathological, but non-malignant, conditions such as hemangiomas, including infantile hemaginomas, angiofibroma of the nasopharynx and avascular necrosis of bone; and disorders of the female reproductive system such as endometriosis. The compounds are also useful for the treatment of edema, and conditions of vascular hyperpermeability.

The compounds of the invention are useful in therapy of proliferative diseases. These compounds can be used for the treatment of an inflammatory rheumatoid or rheumatic disease, especially of manifestations at the locomotor apparatus, such as various inflammatory rheumatoid diseases, especially chronic polyarthritis including rheumatoid arthritis, juvenile arthritis or psoriasis arthropathy; paraneoplastic syndrome or tumor-induced inflammatory diseases, turbid effusions, collagenosis, such as systemic Lupus erythematosus, poly-myositis, dermato-myositis, systemic sclerodermia or mixed collagenosis; postinfectious arthritis (where no living pathogenic organism can be found at or in the affected part of the body), seronegative spondylarthritis, such as spondylitis ankylosans: vasculitis, sarcoidosis, or arthrosis; or further any combinations thereof. An example of an inflammation related disorder is (a) synovial inflammation, for example, synovitis, including any of the particular forms of synovitis, in particular bursal synovitis and purulent symovitis, as far as it is not crystal-induced. Such synovial inflammation may for example, be consequential to or associated with disease, e.g. arthritis, e.g. osteoarthritis, rheumatoid arthritis or arthritis deformans. The present invention is further applicable to the systemic treatment of inflammation, e.g. inflammatory diseases or conditions, of the joints or locomotor apparatus in the

region of the tendon insertions and tendon sheaths. Such

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inflammation may be, for example, consequential to or associated with disease or further (in a broader sense of the invention) with surgical intervention, including, in particular conditions such as insertion endopathy, myofasciale syndrome and tendomyosis. The present invention is further especially applicable to the treatment of

is further especially applicable to the treatment of inflammation, e.g. inflammatory disease or condition, of connective tissues including dermatomyositis and myositis.

These compounds can be used as active agents against such disease states as arthritis, atherosclerosis, psoriasis, hemangiomas, myocardial angiogenesis, coronary and cerebral collaterals, ischemic limb angiogenesis, wound healing, peptic ulcer Helicobacter related diseases, fractures, cat scratch fever, rubeosis, neovascular glaucoma and retinopathies such as those associated with diabetic retinopathy or macular degeneration. In addition, some of these compounds can be used as active agents against solid tumors, malignant ascites, hematopoietic cancers and hyperproliferative disorders such as thyroid hyperplasia (especially Grave's disease), and cysts (such as hypervascularity of ovarian stroma, characteristic of polycystic ovarian syndrome (Stein- Leventhal syndrome)) since such diseases require a proliferation of blood vessel cells for growth and/or metastasis.

Further, some of these compounds can be used as active agents against burns, chronic lung disease, stroke, polyps, anaphylaxis, chronic and allergic inflammation, ovarian hyperstimulation syndrome, brain tumor-associated cerebral edema, high-altitude, trauma or hypoxia induced cerebral or pulmonary edema, ocular and macular edema, ascites, and other diseases where vascular hyperpermeability, effusions, exudates, protein extravasation, or edema is a manifestation of the disease. The compounds will also be useful in treating disorders in which protein extravasation leads to

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the deposition of fibrin and extracellular matrix, promoting stromal proliferation (e.g. fibrosis, cirrhosis and carpal tunnel syndrome).

The compounds of the present invention are also useful

in the treatment of ulcers including bacterial, fungal,

Mooren ulcers and ulcerative colitis.

The compounds of the present invention are also useful

in the treatment of conditions wherein undesired angiogenesis, edema, or stromal deposition occurs in viral infections such as Herpes simplex, Herpes Zoster, AIDS, Kaposi's sarcoma, protozoan infections and toxoplasmosis, following trauma, radiation, stroke, endometriosis, ovarian hyperstimulation syndrome, systemic lupus, sarcoidosis, synovitis, Crohn's disease, sickle cell anaemia, Lyme disease, pemphigoid, Paget's disease, hyperviscosity

disease, pemphigoid, Paget's disease, hyperviscosity syndrome, Osler-Weber-Rendu disease, chronic inflammation, chronic occlusive pulmonary disease, asthma, and inflammatory rheumatoid or rheumatic disease. The compounds are also useful in the reduction of sub-cutaneous fat and

20 for the treatment of obesity.

The compounds of the present invention are also useful in the treatment of ocular conditions such as ocular and macular edema, ocular neovascular disease, scleritis, radial keratotomy, uveitis, vitritis, myopia, optic pits, chronic retinal detachment, post-laser complications, glaucoma, conjunctivitis, Stargardt's disease and Eales disease in addition to retinopathy and macular degeneration.

The compounds of the present invention are also useful in the treatment of cardiovascular conditions such as 30 atherosclerosis, restenosis, arteriosclerosis, vascular occlusion and carotid obstructive disease.

The compounds of the present invention are also useful in the treatment of cancer related indications such as solid tumors, sarcomas (especially Ewing's sarcoma and

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osteosarcoma), retinoblastoma, rhabdomyosarcomas, neuroblastoma, hematopoietic malignancies, including leukemia and lymphoma, tumor- induced pleural or pericardial effusions, and malignant ascites.

5 The compounds of the present invention are also useful in the treatment of diabetic conditions such as diabetic retinopathy and microangiopathy.

The compounds of this invention may also act as inhibitors of other protein kinases, e.g. p38, EGFR, CDK-2, CDK-5, IKK, JNK3, and thus be effective in the treatment of diseases associated with other protein kinases.

Besides being useful for human treatment, these compounds are also useful for veterinary treatment of companion animals, exotic animals and farm animals, including mammals, rodents, and the like. More preferred animals include horses, dogs, and cats.

As used herein, the compounds of the present invention include the pharmaceutically acceptable derivatives thereof.

Definitions

The term "treatment" includes therapeutic treatment as well as prophylactic treatment (either preventing the onset of disorders altogether or delaying the onset of a preclinically evident stage of disorders in individuals).

The term "prevention" includes either preventing the onset of disorders altogether or delaying the onset of a preclinically evident stage of disorders in individuals. This includes prophylactic treatment of those at risk of

30 prophylactic treatment of those at risk of developing a disease, such as a cancer, for example. "Prophylaxis" is another term for prevention. COURSE LOTE OF THE

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A "pharmaceutically-acceptable derivative " denotes any salt, ester of a compound of this invention, or any other compound which upon administration to a patient is capable of providing (directly or indirectly) a compound of this invention, or a metabolite or residue thereof, characterized by the ability to inhibit angiogenesis.

The phrase "therapeutically-effective" is intended to qualify the amount of each agent, which will achieve the goal of improvement in disorder severity and the frequency of incidence over treatment of each agent by itself, while avoiding adverse side effects typically associated with alternative therapies. For example, effective neoplastic therapeutic agents prolong the survivability of the patient, inhibit the rapidly-proliferating cell growth associated with the neoplasm, or effect a regression of the neoplasm.

The term "H" denotes a single hydrogen atom. This radical may be attached, for example, to an oxygen atom to form a hydroxyl radical.

Where the term "alkyl" is used, either alone or within other terms such as "haloalkyl" and "alkylamino", it embraces linear or branched radicals having one to about twelve carbon atoms. More preferred alkyl radicals are "lower alkyl" radicals having one to about six carbon atoms. Examples of such radicals include methyl, ethyl, n-propyl, isopropyl, n-butyl, isobutyl, sec-butyl, tert-butyl, pentyl, isoamyl, hexyl and the like. Even more preferred are lower alkyl radicals having one or two carbon atoms. The term "alkylenyl" embraces bridging divalent alkyl radicals such as methylenyl and ethylenyl. The term "lower alkyl substituted with R2" does not include an acetal moiety.

The term "alkenyl" embraces linear or branched radicals having at least one carbon-carbon double bond of two to about twelve carbon atoms. More preferred alkenyl radicals are "lower alkenyl" radicals having two to about

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six carbon atoms. Most preferred lower alkenyl radicals are radicals having two to about four carbon atoms. Examples of alkenyl radicals include ethenyl, propenyl, allyl, propenyl, butenyl and 4-methylbutenyl. The terms "alkenyl" and "lower alkenyl", embrace radicals having "cis" and "trans" orientations, or alternatively, "E" and "Z" orientations.

The term "alkynyl" denotes linear or branched radicals having at least one carbon-carbon triple bond and having two to about twelve carbon atoms. More preferred alkynyl radicals are "lower alkynyl" radicals having two to about six carbon atoms. Most preferred are lower alkynyl radicals having two to about four carbon atoms. Examples of such radicals include propargyl, butynyl, and the like.

The term "haloalkyl" embraces radicals wherein any one or more of the alkyl carbon atoms is substituted with halo

The term "halo" means halogens such as fluorine, chlorine, bromine or iodine atoms.

as defined above. Specifically embraced are monohaloalkyl, dihaloalkyl and polyhaloalkyl radicals including perhaloalkyl. A monohaloalkyl radical, for one example, may have either an iodo, bromo, chloro or fluoro atom within the radical. Dihalo and polyhaloalkyl radicals may have two or more of the same halo atoms or a combination of different halo radicals. "Lower haloalkyl" embraces radicals having 1-6 carbon atoms. Even more preferred are lower haloalkyl radicals having one to three carbon atoms. Examples of haloalkyl radicals include fluoromethyl, difluoromethyl, trifluoromethyl, chloromethyl, dichloromethyl, trichloromethyl, pentafluoroethyl, heptafluoropropyl, difluorochloromethyl, dichlorofluoromethyl, difluoroethyl, difluoropropyl, dichloroethyl and dichloropropyl.

"Perfluoroalkyl" means alkyl radicals having all hydrogen atoms replaced with fluoro atoms. Examples include

trifluoromethyl and pentafluoroethyl.

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substituent.

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The term "hydroxyalkyl" embraces linear or branched alkyl radicals having one to about ten carbon atoms any one of which may be substituted with one or more hydroxyl radicals. More preferred hydroxyalkyl radicals are "lower hydroxyalkyl" radicals having one to six carbon atoms and one or more hydroxyl radicals. Examples of such radicals include hydroxymethyl, hydroxyethyl, hydroxypropyl, hydroxybutyl and hydroxyhexyl. Even more preferred are lower hydroxyalkyl radicals having one to three carbon atoms.

containing radicals each having alkyl portions of one to about ten carbon atoms. More preferred alkoxy radicals are "lower alkoxy" radicals having one to six carbon atoms. Examples of such radicals include methoxy, ethoxy, propoxy,

The term "alkoxy" embrace linear or branched oxy-

butoxy and tert-butoxy. Even more preferred are lower alkoxy radicals having one to three carbon atoms. Alkoxy radicals may be further substituted with one or more halo atoms, such as fluoro, chloro or bromo, to provide "haloalkoxy" radicals. Even more preferred are lower haloalkoxy radicals having one to three carbon atoms. Examples of such radicals include fluoromethoxy, chloromethoxy, trifluoromethoxy,

trifluoroethoxy, fluoroethoxy and fluoropropoxy.

The term "aryl", alone or in combination, means a carbocyclic aromatic system containing one or two rings wherein such rings may be attached together in a fused manner. The term "aryl" embraces aromatic radicals such as phenyl, naphthyl, indenyl, tetrahydronaphthyl, and indanyl. More preferred aryl is phenyl. Said "aryl" group may have 1 to 3 substituents such as lower alkyl, hydroxyl, halo, haloalkyl, nitro, cyano, alkoxy and lower alkylamino. Phenyl substituted with -O-CH₂-O- forms the aryl benzodioxolyl

The term " heterocyclyl" embraces saturated, partially saturated and unsaturated heteroatom-containing ring

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radicals, where the heteroatoms may be selected from nitrogen, sulfur and oxygen. It does not include rings containing -O-O-,-O-S- or -S-S- portions. Said "heterocyclyl" group may have 1 to 3 substituents such as hydroxyl, Boc, halo, haloalkyl, cyano, lower alkyl, lower aralkyl, oxo, lower alkoxy, amino and lower alkylamino.

Examples of saturated heterocyclic radicals include saturated 3 to 6-membered heteromonocyclic groups containing 1 to 4 nitrogen atoms [e.g. pyrrolidinyl, imidazolidinyl, 10 piperidinyl, pyrrolinyl, piperazinyl]; saturated 3 to 6-membered heteromonocyclic group containing 1 to 2 oxygen atoms and 1 to 3 nitrogen atoms [e.g. morpholinyl]; saturated 3 to 6-membered heteromonocyclic group containing 1 to 2 sulfur atoms and 1 to 3 nitrogen atoms [e.g., 15 thiazolidinyl]. Examples of partially saturated heterocyclyl radicals include dihydrothienyl, dihydropyranyl, dihydrofuryl and dihydrothiezolyl.

Examples of unsaturated heterocyclic radicals, also termed "heteroarvl" radicals, include unsaturated 5 to 6 20 membered heteromonocyclyl group containing 1 to 4 nitrogen atoms, for example, pyrrolyl, imidazolyl, pyrazolyl, 2pyridyl, 3-pyridyl, 4-pyridyl, pyrimidyl, pyrazinyl, pyridazinyl, triazolyl [e.g., 4H-1,2,4-triazolyl, 1H-1,2,3triazolyl, 2H-1,2,3-triazolyl); unsaturated 5- to 6-membered 25 heteromonocyclic group containing an oxygen atom, for example, pyranyl, 2-furyl, 3-furyl, etc.; unsaturated 5 to 6-membered heteromonocyclic group containing a sulfur atom, for example, 2-thienyl, 3-thienyl, etc.; unsaturated 5- to 6-membered heteromonocyclic group containing 1 to 2 oxygen 30 atoms and 1 to 3 nitrogen atoms, for example, oxazolvl, isoxazolyl, oxadiazolyl [e.g., 1,2,4-oxadiazolyl, 1,3,4oxadiazolyl, 1,2,5-oxadiazolyll; unsaturated 5 to 6-membered heteromonocyclic group containing 1 to 2 sulfur atoms and 1 to 3 nitrogen atoms, for example, thiazolyl, thiadiazolyl

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[e.g., 1,2,4-thiadiazolyl, 1,3,4-thiadiazolyl, 1,2,5-thiadiazolyl].

The term also embraces radicals where heterocyclic radicals are fused/condensed with aryl radicals: unsaturated condensed heterocyclic group containing 1 to 5 nitrogen atoms, for example, indolyl, isoindolyl, indolizinyl, benzimidazolyl, quinolyl, isoquinolyl, indazolyl, benzotriazolyl, tetrazolopyridazinyl [e.g.,

tetrazolo [1,5-b]pyridazinyl]; unsaturated condensed

10 heterocyclic group containing 1 to 2 oxygen atoms and 1 to 3 nitrogen atoms [e.g. benzoxazolyl, benzoxadiazolyl]; unsaturated condensed heterocyclic group containing 1 to 2 sulfur atoms and 1 to 3 nitrogen atoms [e.g., benzothiazolyl, benzothiadiazolyl]; and saturated, partially

unsaturated and unsaturated condensed heterocyclic group containing 1 to 2 oxygen or sulfur atoms [e.g. benzofury], benzothieny], 2,3-dihydro-benzo[1,4]dioxinyl and dihydrobenzofury]]. Preferred heterocyclic radicals include five to ten membered fused or unfused radicals. More

20 preferred examples of heteroaryl radicals include quinolyl, isoquinolyl, imidazolyl, pyridyl, thienyl, thiazolyl, oxazolyl, furyl, and pyrazinyl. Other preferred heteroaryl radicals are 5- or 6-membered heteroaryl, containing one or two heteroatoms selected from sulfur, nitrogen and oxygen,

25 selected from thienyl, furyl, pyrrolyl, indazolyl, pyrazolyl, oxazolyl, triazolyl, imidazolyl, pyrazolyl, isoxazolyl, isothiazolyl, pyridyl, piperidinyl and pyrazinyl.

Particular examples of non-nitrogen containing

30 heteroaryl include pyranyl, 2-furyl, 3-furyl, 2-thienyl, 3thienyl, benzofuryl, benzothienyl, and the like.

Particular examples of partially saturated and saturated heterocyclyl include pyrrolidinyl, imidazolidinyl, piperidinyl, pyrrolinyl, pyrazolidinyl, piperazinyl,

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morpholinyl, tetrahydropyranyl, thiazolidinyl, dihydrothienyl, 2,3-dihydro-benzo[1,4]dioxanyl, indolinyl, isoindolinyl, dihydrobenzothienyl, dihydrobenzofuryl, isochromanyl, chromanyl, 1,2-dihydroquinolyl, 1,2,3,4-tetrahydro-isoquinolyl, 1,2,3,4-tetrahydro-quinolyl, 2,3,4,4a,9,9a-hexahydro-1H-3-aza-fluorenyl, 5,6,7-trihydro-1,2,4-triazolo[3,4-a]isoquinolyl, 3,4-dihydro-2H-

1,2,4-triazolo[3,4-a]isoquinolyl, 3,4-dihydro-2H-benzo[1,4]oxazinyl, benzo[1,4]dioxanyl, 2,3-dihydro-1H-1\(\lambda\)'-benzo[d]isothiazol-6-yl, dihydropyranyl, dihydrofuryl and dihydrothiazolyl, and the like.

The term "sulfonyl", whether used alone or linked to other terms such as alkylsulfonyl, denotes respectively divalent radicals -SO₂-.

The terms "sulfamyl," "aminosulfonyl" and "sulfonamidyl," denotes a sulfonyl radical substituted with an amine radical, forming a sulfonamide (-SO:NH₂).

The term "alkylaminosulfonyl" includes "N-alkylaminosulfonyl" where sulfamyl radicals are independently substituted with one or two alkyl radical(s). More preferred alkylaminosulfonyl radicals are "lower alkylaminosulfonyl" radicals having one to six carbon atoms. Even more preferred are lower alkylaminosulfonyl radicals having one to three carbon atoms. Examples of such lower alkylaminosulfonyl radicals include N-methylaminosulfonyl, and N-ethylaminosulfonyl.

The terms "carboxy" or "carboxyl", whether used alone or with other terms, such as "carboxyalkyl", denotes -CO₂H.

The term "carbonyl", whether used alone or with other terms, such as "aminocarbonyl", denotes -(C=0)-.

30 The term "aminocarbonyl" denotes an amide group of the formula -C(=0)NH₂.

The terms "N-alkylaminocarbonyl" and "N,N-dialkylaminocarbonyl" denote aminocarbonyl radicals independently substituted with one or two alkyl radicals,

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respectively. More preferred are "lower alkylaminocarbonyl" having lower alkyl radicals as described above attached to an aminocarbonyl radical.

The terms "N-arylaminocarbonyl" and "N-alkyl-N-arylaminocarbonyl" denote aminocarbonyl radicals substituted, respectively, with one aryl radical, or one alkyl and one aryl radical.

The term "heterocyclylalkylenyl" embraces heterocyclic-substituted alkyl radicals. More preferred heterocyclylalkylenyl radicals are "5- or 6-membered heteroarylalkylenyl" radicals having alkyl portions of one to six carbon atoms and a 5- or 6-membered heteroaryl radical. Even more preferred are lower heteroarylalkylenyl radicals having alkyl portions of one to three carbon atoms. Examples include such radicals as pyridylmethyl and thienylmethyl.

The term "aralkyl" embraces aryl-substituted alkyl radicals. Preferable aralkyl radicals are "lower aralkyl" radicals having aryl radicals attached to alkyl radicals having one to six carbon atoms. Even more preferred are "phenylalkylenyl" attached to alkyl portions having one to three carbon atoms. Examples of such radicals include benzyl, diphenylmethyl and phenylethyl. The aryl in said aralkyl may be additionally substituted with halo, alkyl, alkoxy, halkoalkyl and haloalkoxy.

The term "alkylthio" embraces radicals containing a linear or branched alkyl radical, of one to ten carbon atoms, attached to a divalent sulfur atom. Even more preferred are lower alkylthio radicals having one to three carbon atoms. An example of "alkylthio" is methylthio, (CH₁S-).

The term "haloalkylthio" embraces radicals containing a haloalkyl radical, of one to ten carbon atoms, attached to a divalent sulfur atom. Even more preferred are lower

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haloalkylthio radicals having one to three carbon atoms. An example of "haloalkylthio" is trifluoromethylthio.

The term "alkylamino" embraces "N-alkylamino" and "N,N-dialkylamino" where amino groups are substituted with one alkyl radical and with two independent alkyl radicals, respectively. More preferred alkylamino radicals are "lower alkylamino" radicals having one or two alkyl radicals of one to six carbon atoms, attached to a nitrogen atom. Even more preferred are lower alkylamino radicals having one to three carbon atoms. Suitable alkylamino radicals may be mono or dialkylamino such as N-methylamino, N-ethylamino, N,N-diethylamino and the like.

The term "arylamino" denotes amino groups which have been substituted with one or two aryl radicals, such as Nphenylamino. The arylamino radicals may be further substituted on the aryl ring portion of the radical.

The term "heteroarylamino" denotes amino groups which have been substituted with one or two heteroaryl radicals, such as N-thienylamino. The "heteroarylamino" radicals may be further substituted on the heteroaryl ring portion of the radical.

The term "aralkylamino" denotes amino groups which have been substituted with one or two aralkyl radicals. More preferred are phenyl-C₁-C₃-alkylamino radicals, such as N-benzylamino. The aralkylamino radicals may be further substituted on the aryl ring portion.

The terms "N-alkyl-N-arylamino" and "N-aralkyl-N-alkylamino" denote amino groups which have been substituted with one aralkyl and one alkyl radical, or one aryl and one alkyl radical, respectively, to an amino group.

The term "aminoalkyl" embraces linear or branched alkyl radicals having one to about ten carbon atoms any one of which may be substituted with one or more amino radicals. More preferred aminoalkyl radicals are "lower aminoalkyl"

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radicals having one to six carbon atoms and one or more amino radicals. Examples of such radicals include aminomethyl, aminoethyl, aminopropyl, aminobutyl and aminohexyl. Even more preferred are lower aminoalkyl radicals having one to three carbon atoms.

The term "alkylaminoalkyl" embraces alkyl radicals substituted with alkylamino radicals. More preferred alkylaminoalkyl radicals are "lower alkylaminoalkyl" radicals having alkyl radicals of one to six carbon atoms. Even more preferred are lower alkylaminoalkyl radicals having alkyl radicals of one to three carbon atoms. Suitable alkylaminoalkyl radicals may be mono or dialkyl substituted, such as N-methylaminomethyl, N,N-dimethyl-aminoethyl, N,N-diethylaminomethyl and the like.

The term "alkylaminoalkoxy" embraces alkoxy radicals substituted with alkylamino radicals. More preferred alkylaminoalkoxy radicals are "lower alkylaminoalkoxy" radicals having alkoxy radicals of one to six carbon atoms. Even more preferred are lower alkylaminoalkoxy radicals having alkyl radicals of one to three carbon atoms. Suitable alkylaminoalkoxy radicals may be mono or dialkyl substituted, such as N-methylaminoethoxy, N,N-dimethylaminoethoxy, N,N-dimethylaminoethoxy, and the like.

The term "alkylaminoalkoxyalkoxy" embraces alkoxy

25 radicals substituted with alkylaminoalkoxy radicals. More
preferred alkylaminoalkoxyalkoxy radicals are "lower
alkylaminoalkoxyalkoxy" radicals having alkoxy radicals of
one to six carbon atoms. Even more preferred are lower
alkylaminoalkoxyalkoxy radicals having alkyl radicals of one
30 to three carbon atoms. Suitable alkylaminoalkoxyalkoxy
radicals may be mono or dialkyl substituted, such as Nmethylaminoethoxyethoxy, N,N-dimethylaminoethoxyethoxy, N,Ndiethylaminomethoxymethoxy and the like.

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The term "carboxyalkyl" embraces linear or branched alkyl radicals having one to about ten carbon atoms any one of which may be substituted with one or more carboxy radicals. More preferred carboxyalkyl radicals are "lower carboxyalkyl" radicals having one to six carbon atoms and one carboxy radical. Examples of such radicals include carboxymethyl, carboxypropyl, and the like. Even more preferred are lower carboxyalkyl radicals having one to three CH₃ groups.

The term "halosulfonyl" embraces sulfonyl radicals substituted with a halogen radical. Examples of such halosulfonyl radicals include chlorosulfonyl and fluorosulfonyl.

The term "arylthio" embraces aryl radicals of six to ten carbon atoms, attached to a divalent sulfur atom. An example of "arylthio" is phenylthio.

The term "aralkylthio" embraces aralkyl radicals as described above, attached to a divalent sulfur atom. More preferred are phenyl- C_1 - C_3 -alkylthio radicals. An example of "aralkylthio" is benzylthio.

The term "aryloxy" embraces optionally substituted aryl radicals, as defined above, attached to an oxygen atom. Examples of such radicals include phenoxy.

The term "aralkoxy" embraces oxy-containing aralkyl radicals attached through an oxygen atom to other radicals. More preferred aralkoxy radicals are "lower aralkoxy" radicals having optionally substituted phenyl radicals attached to lower alkoxy radical as described above.

The term "heteroaryloxy" embraces optionally substituted heteroaryl radicals, as defined above, attached to an oxygen atom.

The term "heteroarylalkoxy" embraces oxy-containing heteroarylalkyl radicals attached through an oxygen atom to other radicals. More preferred heteroarylalkoxy radicals are

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"lower heteroarylalkoxy" radicals having optionally substituted heteroaryl radicals attached to lower alkoxy radical as described above.

The term "cycloalkyl" includes saturated carbocyclic groups. Preferred cycloalkyl groups include C_3 - C_6 rings. More preferred compounds include, cyclopentyl, cyclopropyl, and cyclohexyl.

The term "cycloalkenyl" includes carbocyclic groups having one or more carbon-carbon double bonds including "cycloalkyldienyl" compounds. Preferred cycloalkenyl groups include C₃-C₆ rings. More preferred compounds include, for example, cyclopentenyl, cyclopentadienyl, cyclohexenyl and cycloheptadienyl.

The term "comprising" is meant to be open ended,

15 including the indicated component but not excluding other
elements.

The phrase "Formula I-XII" includes sub formulas such as II'.

The compounds of the invention are endowed with kinase inhibitory activity, such as KDR inhibitory activity.

The present invention also comprises the use of a compound of the invention, or pharmaceutically acceptable salt thereof, in the manufacture of a medicament for the treatment either acutely or chronically of an angiogenesis mediated disease state, including those described previously. The compounds of the present invention are useful in the manufacture of an anti-cancer medicament. The compounds of the present invention are also useful in the manufacture of a medicament to attenuate or prevent disorders through inhibition of KDR.

The present invention comprises a pharmaceutical composition comprising a therapeutically-effective amount of a compound of Formulas I-XII in association with a least one pharmaceutically-acceptable carrier, adjuvant or diluent.

The present invention also comprises a method of treating angiogenesis related disorders in a subject having or susceptible to such disorder, the method comprising treating the subject with a therapeutically-effective amount of a compound of Formula I

wherein each of A^1 and A^2 is independently C, CH or N; 10 wherein ring A is selected from

a) 5- or 6-membered partially saturated heterocyclyl,

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- b) 5- or 6-membered heteroarvl,
- c) 9-, 10- or 11-membered fused partially saturated heterocycly1,
- d) 9-, 10- or 11-membered fused heteroarvl;
- e) naphthyl, and
- f) 4-, 5- or 6- membered cycloalkenyl;

wherein X is

wherein Z is oxygen or sulfur;

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wherein p is 0 to 2,

wherein R^a and R^b are independently selected from H, halo,

cyano, -NHR⁶ and C_{1-4} -alkyl substituted with R^2 , or wherein R^a and R^b together form C_3 - C_6 cycloalkyl;

wherein R^z is selected from C_2 - C_6 -alkylenyl, where one of the CH₂ groups may be replaced with an oxygen atom or an -NH-; wherein one of the CH₂ groups may be substituted with one or two radicals selected from halo, cyano, -NHR⁶ and C_{1-4} -alkyl substituted with R^2 ;

wherein R^d is cycloalkyl;

wherein R is selected from

- a) substituted or unsubstituted 5-6 membered heterocyclyl, b) substituted aryl, and
- c) substituted or unsubstituted fused 9-14-membered bicyclic or tricyclic heterocyclyl; wherein substituted R is substituted with one or more substituents independently selected from halo, -OR³, -SR², -SO₂R³, -CO₂R³, -COR³, -NR³R³, -SO₂NR³R³, -SO₂NR³R³, -NR²C(O)OR³, -NR²C(O)R³, cycloalkyl, optionally substituted 5-6 membered heterocyclyl, optionally substituted phenyl, nitro, alkylaminoalkoxyalkoxy, cyano, alkylaminoalkoxy, lower alkyl substituted with R², lower alkenyl substituted with R², and lower alkynyl substituted with R²,

wherein R1 is selected from

- a) substituted or unsubstituted 6-10 membered aryl,
- b) substituted or unsubstituted 5-6 membered 30 heterocycly1,

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- c) substituted or unsubstituted 9-14 membered bicyclic or tricyclic heterocyclyl,
- d) cycloalkyl, and
- e) cycloalkenyl,
- 5 wherein substituted R¹ is substituted with one or more substituents independently selected from halo, -OR³, -SR³, -CO₂R³, -CONR³R³, -COR³, -NR³R³, -NR (C₁-C₄ alkylenylR³⁴), -SO₂R³, -SO₂NR³R³, -NR³C(O)OR³, -NR³C(O)R³, optionally substituted cycloalkyl, optionally substituted 5-6 membered heterocyclyl, optionally substituted phenyl, halosulfonyl, cyano, alkylaminoalkoxy, alkylaminoalkoxyalkoxy, nitro, lower alkyl substituted with R², lower alkenyl substituted with R², and lower alkynyl substituted with R²:
 - wherein R² is one or more substituents independently selected from H, halo, -OR³, oxo, -SR³, -CO₂R³, -COR³, -COR³, -CORR³R³, -NR³R³, -SO₂NR³R³, -NR³C(O)OR³, -NR³C(O)R³, cycloalkyl, optionally substituted phenylalkylenyl, optionally substituted 5-6 membered heterocyclyl, optionally substituted heteroarylalkylenyl, optionally substituted phenyl, lower alkyl, cyano, lower hydroxyalkyl, lower carboxyalkyl, nitro, lower alkenyl, lower alkynyl, lower

aminoalkyl, lower alkylaminoalkyl and lower haloalkyl;

- 25 wherein R³ is selected from H, lower alkyl, phenyl, heterocyclyl, C3-C6-cycloalkyl, phenylalkyl, heterocyclylalkyl, C3-C6 cycloalkylalkyl, and lower haloalkyl;
- wherein R^4 is selected from a direct bond, C_{2-4} -alkylenyl, C_{2-3} and C_{2-4} -alkynylenyl, where one of the CH_2 groups may be substituted with an oxygen atom or an -NH-, wherein R^4 is optionally substituted with hydroxy;
 - wherein R⁵ is selected from H, lower alkyl, phenyl and lower aralkyl;

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wherein R^{5a} is selected from H, lower alkyl, phenyl and lower aralkyl:

wherein R^6 is selected from H or C_{1-6} -alkyl; and wherein R^{14} is selected from H, phenyl, 5-6 membered

heterocyclyl and C_3 - C_6 cycloalkyl; and pharmaceutically acceptable derivatives thereof; provided A is not naphthyl when X is -C(0)NH- and when R¹ is phenyl when Y is -NCH₂- and when R is 4-pyridyl; and further provided R is not unsubstituted 2-thienyl, 2-pyridyl or 3pyridyl when Y is -NHCH₂-.

COMBINATIONS

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While the compounds of the invention can be administered as the sole active pharmaceutical agent, they can also be used in combination with one or more compounds of the invention or other agents. When administered as a combination, the therapeutic agents can be formulated as separate compositions that are administered at the same time or sequentially at different times, or the therapeutic agents can be given as a single composition.

The phrase "co-therapy" (or "combination-therapy"), in defining use of a compound of the present invention and another pharmaceutical agent, is intended to embrace administration of each agent in a sequential manner in a regimen that will provide beneficial effects of the drug combination, and is intended as well to embrace co-administration of these agents in a substantially simultaneous manner, such as in a single capsule having a fixed ratio of these active agents or in multiple, separate capsules for each agent.

Specifically, the administration of compounds of the present invention may be in conjunction with additional therapies known to those skilled in the art in the

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prevention or treatment of neoplasia, such as with radiation therapy or with cytostatic or cytotoxic agents.

If formulated as a fixed dose, such combination products employ the compounds of this invention within the accepted dosage ranges. Compounds of Formula I may also be administered sequentially with known anticancer or cytotoxic agents when a combination formulation is inappropriate. The invention is not limited in the sequence of administration; compounds of the invention may be administered either prior to, simultaneous with, or after administration of the known anticancer or cytotoxic agent.

Currently, standard treatment of primary tumors consists of surgical excision followed by either radiation or IV administered chemotherapy. The typical chemotherapy regime consists of either DNA alkylating agents, DNA intercalating agents, CDK inhibitors, or microtubule poisons. The chemotherapy doses used are just below the maximal tolerated dose and therefore dose limiting toxicities typically include, nausea, vomiting, diarrhea, hair loss, neutropenia and the like.

There are large numbers of antineoplastic agents available in commercial use, in clinical evaluation and in pre-clinical development, which would be selected for treatment of neoplasia by combination drug chemotherapy. Such antineoplastic agents fall into several major

categories, namely, antibiotic-type agents, alkylating agents, antimetabolite agents, hormonal agents, immunological agents, interferon-type agents and a category of miscellaneous agents.

30 A first family of antineoplastic agents which may be used in combination with compounds of the present invention consists of antimetabolite-type/thymidilate synthase inhibitor antineoplastic agents. Suitable antimetabolite antineoplastic agents may be selected from but not limited

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to the group consisting of 5-FU-fibrinogen, acanthifolic acid, aminothiadiazole, brequinar sodium, carmofur, Ciba-Geigy CGP-30694, cyclopentyl cytosine, cytarabine phosphate stearate, cytarabine conjugates, Lilly DATHF, Merrel Dow

- DDFC, dezaguanine, dideoxycytidine, dideoxyguanosine, didox, Yoshitomi DMDC, doxifluridine, Wellcome EHNA, Merck & Co. EX-015, fazarabine, floxuridine, fludarabine phosphate, 5fluorouracil, N-(2'-furanidyl)-5-fluorouracil, Daiichi Seiyaku FO-152, isopropyl pyrrolizine, Lilly LY-188011,
- Lilly LY-264618, methobenzaprim, methotrexate, Wellcome MZPES, norspermidine, NCI NSC-127716, NCI NSC-264880, NCI NSC-39661, NCI NSC-612567, Warner-Lambert PALA, pentostatin, piritrexim, plicamycin, Asahi Chemical PL-AC, Takeda TAC-788, thioguanine, tiazofurin, Erbamont TIF, trimetrexate, tyrosine kinase inhibitors, Taiho UFT and uricytin.

A second family of antineoplastic agents which may be used in combination with compounds of the present invention consists of alkylating-type antineoplastic agents. Suitable alkylating-type antineoplastic agents may be selected from but not limited to the group consisting of Shionogi 254-S, aldo-phosphamide analogues, altretamine, anaxirone, Boehringer Mannheim BBR-2207, bestrabucil, budotitane, Wakunaga CA-102, carboplatin, carmustine, Chinoin-139, Chinoin-153, chlorambucil, cisplatin, cyclophosphamide,

- American Cyanamid CL-286558, Sanofi CY-233, cyplatate, Degussa D-19-384, Sumimoto DACHP(Myr)2, diphenylspiromustine, diplatinum cytostatic, Erba distamycin derivatives, Chugai DWA-2114R, ITI E09, elmustine, Erbamont FCE-24517, estramustine phosphate sodium, fotemustine,
- 30 Unimed G-6-M, Chinoin GYKI-17230, hepsul-fam, ifosfamide, iproplatin, lomustine, mafosfamide, mitolactol, Nippon Kayaku NK-121, NCI NSC-264395, NCI NSC-342215, oxaliplatin, Upjohn PCNU, prednimustine, Proter PTT-119, ranimustine, semustine, SmithKline SK&F-101772, Yakult Honsha SN-22,

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spiromus-tine, Tanabe Seiyaku TA-077, tauromustine, temozolomide, teroxirone, tetraplatin and trimelamol.

A third family of antineoplastic agents which may be used in combination with compounds of the present invention consists of antibiotic-type antineoplastic agents. Suitable antibiotic-type antineoplastic agents may be selected from but not limited to the group consisting of Taiho 4181-A, aclarubicin, actinomycin D, actinoplanone, Erbamont ADR-456, aeroplysinin derivative, Ajinomoto AN-201-II, Ajinomoto AN-3, Nippon Soda anisomycins, anthracycline, azino-mycin-A,

- 3, Nippon Soda anisomycins, anthracycline, azino-mycin-A, bisucaberin, Bristol-Myers BL-6859, Bristol-Myers BMY-25067, Bristol-Myers BMY-25551, Bristol-Myers BMY-26605, Bristol-Myers BMY-27557, Bristol-Myers BMY-28438, bleomycin sulfate, bryostatin-1, Taiho C-1027, calichemycin, chromoximycin, dautonywini, dauton
- dactinomycin, daunorubicin, Kyowa Hakko DC-102, Kyowa Hakko DC-79, Kyowa Hakko DC-88A, Kyowa Hakko DC92-B, ditrisarubicin B, Shionogi DOB-41, doxorubicin, doxorubicin-fibrinogen, elsamicin-A, epirubicin, erbstatin, esorubicin, esperamicin-Al, esperamicin-Alb, Erbamont FCE-20 21954, Pulisawa FK-973, fostriecin, Pulisawa FR-900482.
- 20 21954, Fujisawa FK-973, fostriecin, Fujisawa FR-900482, glidobactin, gregatin-A, grincamycin, herbimycin, idarubicin, illudins, kazusamycin, kesarirhodins, Kyowa Hakko KM-5539, Kirin Brewery KRN-8602, Kyowa Hakko KT-5432, Kyowa Hakko KT-5594, Kyowa Hakko KT-6149, American Cyanamid
- 25 LL-D49194, Meiji Seika ME 2303, menogaril, mitomycin, mitoxantrone, SmithKline M-TAG, neoenactin, Nippon Kayaku NK-313, Nippon Kayaku NKT-01, SRI International NSC-357704, oxalysine, oxaunomycin, peplomycin, pilatin, pirarubicin, porothramycin, pyrindanycin A, Tobishi RA-I, rapamycin,
- 30 rhizoxin, rodorubicin, sibanomicin, siwenmycin, Sumitomo SM-5887, Snow Brand SN-706, Snow Brand SN-07, sorangicin-A, sparsomycin, SS Pharmaceutical SS-21020, SS Pharmaceutical SS-7313B, SS Pharmaceutical SS-9816B, steffimycin B, Taiho 4181-2, talisomycin, Takeda TAN-868A, terpentecin, thrazine,

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tricrozarin A, Upjohn U-73975, Kyowa Hakko UCN-10028A, Fujisawa WF-3405, Yoshitomi Y-25024 and zorubicin.

A fourth family of antineoplastic agents which may be used in combination with compounds of the present invention consists of a miscellaneous family of antineoplastic agents. including tubulin interacting agents, topoisomerase II inhibitors, topoisomerase I inhibitors and hormonal agents, selected from but not limited to the group consisting of Qcarotene, α-difluoromethyl-arginine, acitretin, Biotec AD-5, Kyorin AHC-52, alstonine, amonafide, amphethinile, amsacrine, Angiostat, ankinomycin, anti-neoplaston A10, antineoplaston A2, antineoplaston A3, antineoplaston A5, antineoplaston AS2-1, Henkel APD, aphidicolin glycinate, asparaginase, Avarol, baccharin, batracylin, benfluron, benzotript, Ipsen-Beaufour BIM-23015, bisantrene, Bristol-Myers BMY-40481, Vestar boron-10, bromofosfamide, Wellcome BW-502, Wellcome BW-773, caracemide, carmethizole hydrochloride, Ajinomoto CDAF, chlorsulfaquinoxalone, Chemes CHX-2053, Chemex CHX-100, Warner-Lambert CI-921, Warner-Lambert CI-937, Warner-Lambert CI-941, Warner-Lambert CI-958, clanfenur, claviridenone, ICN compound 1259, ICN compound 4711, Contracan, Yakult Honsha CPT-11, crisnatol, curaderm, cytochalasin B, cytarabine, cytocytin, Merz D-609, DABIS maleate, dacarbazine, datelliptinium, didemnin-B, dihaematoporphyrin ether, dihydrolenperone, dinaline, distamycin, Toyo Pharmar DM-341, Toyo Pharmar DM-75, Daiichi Seiyaku DN-9693, docetaxel elliprabin, elliptinium acetate, Tsumura EPMTC, the epothilones, ergotamine, etoposide, etretinate, fenretinide, Fujisawa FR-57704, gallium nitrate, genkwadaphnin, Chugai GLA-43, Glaxo GR-63178, grifolan NMF-5N, hexadecylphosphocholine, Green Cross HO-221,

homoharringtonine, hydroxyurea, BTG ICRF-187, ilmofosine, isoglutamine, isotretinoin, Otsuka JI-36, Ramot K-477, Otsuak K-76COONa, Kureha Chemical K-AM, MECT Corp KI-8110,

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American Cvanamid L-623, leukoregulin, lonidamine, Lundbeck LU-23-112, Lilly LY-186641, NCI (US) MAP, marycin, Merrel Dow MDL-27048, Medco MEDR-340, merbarone, merocyanine derivatives, methylanilinoacridine, Molecular Genetics MGI-136, minactivin, mitonafide, mitoquidone mopidamol, motretinide. Zenyaku Kogyo MST-16. N-(retinovl)amino acids. Nisshin Flour Milling N-021, N-acvlated-dehydroalanines, nafazatrom, Taisho NCU-190, nocodazole derivative, Normosang, NCI NSC-145813, NCI NSC-361456, NCI NSC-604782, NCI NSC-95580, ocreotide, Ono ONO-112, oquizanocine, Akzo Org-10172, paclitaxel, pancratistatin, pazelliptine, Warner-Lambert PD-111707, Warner-Lambert PD-115934, Warner-Lambert PD-131141, Pierre Fabre PE-1001, ICRT peptide D, piroxantrone, polyhaematoporphyrin, polypreic acid, Efamol porphyrin, probimane, procarbazine, proglumide, Invitron protease nexin I, Tobishi RA-700, razoxane, Sapporo Breweries RBS, restrictin-P, retelliptine, retinoic acid, Rhone-Poulenc RP-49532, Rhone-Poulenc RP-56976, SmithKline

SK&F-104864, Sumitomo SM-108, Kuraray SMANCS, SeaPharm SP20 10094, spatol, spirocyclopropane derivatives,
spirogermanium, Unimed, SS Pharmaceutical SS-554,
strypoldinone, Stypoldione, Suntory SUN 0237, Suntory SUN
2071, superoxide dismutase, Toyama T-506, Toyama T-680,
taxol, Teijin TEI-0303, teniposide, thaliblastine, Eastman

25 Kodak TJB-29, tocotrienol, topotecan, Topostin, Teijin TT-82, Kyowa Hakko UCN-01, Kyowa Hakko UCN-1028, ukrain, Eastman Kodak USB-006, vinblastine sulfate, vincristine, vindesine, vinestramide, vinorelbine, vintriptol, vinzolidine, withanolides and Yamanouchi YM-534.

30 Alternatively, the present compounds may also be used in co-therapies with other anti-neoplastic agents, such as acemannan, aclarubicin, aldesleukin, alemtuzumab, alitretinoin, altretamine, amifostine, aminolevulinic acid, amrubicin, amsacrine, anagrelide, anastrozole, ANCER, A-733A - 99 -

ancestim, ARGLABIN, arsenic trioxide, BAM 002 (Novelos), bexarotene, bicalutamide, broxuridine, capecitabine, celmoleukin, cetrorelix, cladribine, clotrimazole, cytarabine ocfosfate, DA 3030 (Dong-A), daclizumab, denileukin diftitox, deslorelin, dexrazoxane, dilazep, docetaxel, docosanol, doxercalciferol, doxifluridine, doxorubicin, bromocriptine, carmustine, cytarabine,

fluorouracil, HIT diclofenac, interferon alfa, daunorubicin, doxorubicin, tretinoin, edelfosine, edrecolomab, eflornithine, emitefur, epirubicin, epoetin

- edrecolomab, eflornithine, emitefur, epirubicin, epoetin beta, etoposide phosphate, exemestane, exisulind, fadrozole, filgrastim, finasteride, fludarabine phosphate, formestane, fotemustine, gallium nitrate, gemcitabine, gemtuzumab zogamicin, gimeracil/oteracil/tegafur
- 15 combination, glycopine, goserelin, heptaplatin, human chorionic gonadotropin, human fetal alpha fetoprotein, ibandronic acid, idarubicin, (imiquimod, interferon alfa, interferon alfa, natural, interferon alfa-2, interferon alfa-2a, interferon alfa-2b, interferon alpha, natural.
 20 alfa-n3, interferon alfacon-1, interferon alpha, natural.
 - alfa-n3, interferon alfacon-1, interferon alpha, natural, interferon beta, interferon beta-1b, interferon gamma, natural interferon gamma-1a, interferon gamma-1b, interleukin-1 beta, iobenguane, irinotecan, irsogladine, lanreotide, LC 9018 (Yakult), leflunomide,
- 25 lenograstim, lentinan sulfate, letrozole, leukocyte alpha interferon, leuprorelin, levamisole + fluorouracil, liarozole, lobaplatin, lonidamine, lovastatin, masoprocol, melarsoprol, metoclopramide, mifepristone, miltefosine, mirimostim, mismatched double stranded RNA, mitoguazone,
- 30 mitolactol, mitoxantrone, molgramostim, nafarelin, naloxone + pentazocine, nartograstim, nedaplatin, nilutamide, noscapine, novel erythropoiesis stimulating protein, NSC 631570 octreotide, oprelvekin, osaterone, oxaliplatin, paclitaxel, pamidronic acid, pegaspargase, peginterferon

alfa-2b, pentosan polysulfate sodium, pentostatin, picibanil, pirarubicin, rabbit antithymocyte polyclonal antibody, polyethylene glycol interferon alfa-2a, porfimer sodium, raloxifene, raltitrexed, rasburicase, rhenium Re

- 186 etidronate, RII retinamide, rituximab, romurtide, samarium (153 Sm) lexidronam, sargramostim, sizofiran, sobuzoxane, sonermin, strontium-89 chloride, suramin, tasonermin, tazarotene, tegafur, temoporfin, temozolomide, teniposide, tetrachlorodecaoxide, thalidomide, thymalfasin,
- thyrotropin alfa, topotecan, toremifene, tositumomab-iodine 131, trastuzumab, treosulfan, tretinoin, trilostane, trimetrexate, triptorelin, tumor necrosis factor alpha, natural, ubenimex, bladder cancer vaccine, Maruyama vaccine, melanoma lysate vaccine, valrubicin, verteporfin.
- vinorelbine, VIRULIZIN, zinostatin stimalamer, or zoledronic acid; abarelix; AE 941 (Aeterna), ambamustine, antisense oligonucleotide, bcl-2 (Genta), APC 8015 (Dendreon), cetuximab, decitabine, dexaminoglutethimide, diaziguone, EL 532 (Elan), EM 800 (Endorecherche).
- 20 eniluracil, etanidazole, fenretinide, filgrastim SD01 (Amgen), fulvestrant, galocitabine, gastrin 17 immunogen, HLA-B7 gene therapy (Vical), granulocyte macrophage colony stimulating factor, histamine dihydrochloride, ibritumomab tiuxetan, ilomastat, IM 862 (Cytran), interleukin-2,
- 25 iproxifene, LDI 200 (Milkhaus), leridistim, lintuzumab, CA 125 MAb (Biomira), cancer MAb (Japan Pharmaceutical Development), HER-2 and Fc MAb (Medarex), idiotypic 105AD7 MAb (CRC Technology), idiotypic CEA MAb (Trilex), LYM-1iodine 131 MAb (Techniclone), polymorphic epithelial mucin-30 yttrium 90 MAb (Antisoma), marimastat, menogaril,
- mitumomab, motexafin gadolinium, MX 6 (Galderma), nelarabine, nolatrexed, P 30 protein, pegvisomant, pemetrexed, porfiromycin, prinomastat, RL 0903 (Shire), rubitecan, satraplatin, sodium phenylacetate, sparfosic

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acid, SRL 172 (SR Pharma), SU 5416 (SUGEN), TA 077 (Tanabe), tetrathiomolybdate, thaliblastine, thrombopoietin, tin ethyl etiopurpurin, tirapazamine, cancer vaccine (Biomira), melanoma vaccine (New York University), melanoma vaccine (Sloan Kettering Institute), melanoma oncolysate vaccine (New York Medical College), viral melanoma cell lysates vaccine (Royal Newcastle Hospital), or valspodar.

Alternatively, the present compounds may also be used in co-therapies with other anti-neoplastic agents, such as other kinase inhibitors including p38 inhibitors and CDK inhibitors, TNF inhibitors, metallomatrix proteases inhibitors (MMP), COX-2 inhibitors including celecoxib, rofecoxib, parecoxib, valdecoxib, and etoricoxib, NSAID's, SOD mimics or α , β 3 inhibitors.

The present invention comprises processes for the preparation of a compound of Formula I-XII.

Also included in the family of compounds of Formula I-XII are the pharmaceutically-acceptable salts thereof. The term "pharmaceutically-acceptable salts" embraces salts commonly used to form alkali metal salts and to form addition salts of free acids or free bases. The nature of the salt is not critical, provided that it is pharmaceutically-acceptable. Suitable pharmaceuticallyacceptable acid addition salts of compounds of Formula I-XII may be prepared from an inorganic acid or from an organic acid. Examples of such inorganic acids are hydrochloric, hydrobromic, hydroiodic, nitric, carbonic, sulfuric and phosphoric acid. Appropriate organic acids may be selected from aliphatic, cycloaliphatic, aromatic, arylaliphatic, heterocyclic, carboxylic and sulfonic classes of organic acids, example of which are formic, acetic, adipic, butyric, propionic, succinic, glycolic, gluconic, lactic, malic, tartaric, citric, ascorbic, glucuronic, maleic, fumaric,

pyruvic, aspartic, glutamic, benzoic, anthranilic, mesylic, 4-hydroxybenzoic, phenylacetic, mandelic, embonic (pamoic), methanesulfonic, ethanesulfonic, ethanedisulfonic, benzenesulfonic, pantothenic, 2-hydroxyethanesulfonic,

- benzenesulfonic, pantochenic, 2-nydroxyethanesulfonic, toluenesulfonic, sulfanilic, cyclohexylaminosulfonic, camphoric, camphorsulfonic, digluconic, cyclopentanepropionic, dodecylsulfonic, glucoheptanoic, glycerophosphonic, heptanoic, hexanoic, 2-hydroxyethanesulfonic, nicotinic, 2-naphthalenesulfonic, oxalic,
- palmoic, pectinic, persulfuric, 2-phenylpropionic, picric, pivalic propionic, succinic, tartaric, thiocyanic, mesylic, undecanoic, stearic, algenic, β-hydroxybutyric, salicylic, galactaric and galacturonic acid. Suitable pharmaceutically-acceptable base addition salts of compounds of Formula I-XII
- include metallic salts, such as salts made from aluminum, calcium, lithium, magnesium, potassium, sodium and zinc, or salts made from organic bases including primary, secondary and tertiary amines, substituted amines including cyclic amines, such as caffeine, arginine, diethylamine, N-ethyl
- 20 piperidine, aistidine, glucamine, isopropylamine, lysine, morpholine, N-ethyl morpholine, piperazine, piperidine, triethylamine, trimethylamine. All of these salts may be prepared by conventional means from the corresponding compound of the invention by reacting, for example, the
- 25 appropriate acid or base with the compound of Formula I-XII.

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Also, the basic nitrogen-containing groups can be quaternized with such agents as lower alkyl halides, such as methyl, ethyl, propyl, and butyl chloride, bromides and iodides; dialkyl sulfates like dimethyl, diethyl, dibutyl, and diamyl sulfates, long chain halides such as decyl, lauryl, myristyl and stearyl chlorides, bromides and iodides, aralkyl halides like benzyl and phenethyl bromides, and others. Water or oil-soluble or dispersible products are thereby obtained.

Examples of acids that may be employed to form pharmaceutically acceptable acid addition salts include such inorganic acids as hydrochloric acid, sulphuric acid and phosphoric acid and such organic acids as oxalic acid, maleic acid, succinic acid and citric acid. Other examples include salts with alkali metals or alkaline earth metals, such as sodium, potassium, calcium or magnesium or with organic bases. Preferred salts include hydrochloride, phosphate and edisylate.

Additional examples of such salts can be found in 20 Berge et al., J. Pharm. Sci., 66, 1 (1977).

GENERAL SYNTHETIC PROCEDURES

The compounds of the invention can be synthesized
25 according to the following procedures of Schemes 1-48,
wherein the substituents are as defined for Formulas I-XII,
above, except where further noted.

Scheme 1

Cyclic amides can be prepared according to the method set out in Scheme 1. The amino group of compound 1 (where R° is alkyl, aryl, and the like) is protected, such as with Boc anhydride, followed by treatment, to remove the ester, 10 such as with base, forming the protected amine/free acid 2. Alternatively, other amino protecting groups known in the art can be used. Substituted amines are coupled with the free acid, such as with EDC, to form the protected amine/amide 3. The protected amine moiety is deprotected, 15 such as with acid, and reacted via one step reductive alkylation with carbonyl-containing compounds (where R' is H, halo, cyano, -NHR6 and C1-4 alkyl) to form the 1-amido-2substituted amino-compounds 4. Preferably the amination is in an alcohol, such as MeOH, EtOH or propanol, and at a temperature between about 0-50°C, such as RT. Aldehydes or 20

ketones are preferred carbonyl-containing compounds.

Alternative carbonyl-containing compounds are, for example, bisulfite adducts or hemiacetals, acetals, hemiketals or ketals of compounds with alcohols, for example lower

- 5 hydroxyalkyl compounds; or thioacetals or thioketals of compounds with mercaptans, for example lower alkylthio compounds. The reductive alkylation is preferably carried out with hydrogenation in the presence of a catalyst, such as platinum or especially palladium, which is preferably
- 10 bonded to a carrier material, such as carbon, or a heavy metal catalyst, such as Raney nickel, at normal pressure or at pressures of from 0.1 to 10 MegaPascal (MPa), or with reduction by means of complex hydrides, such as borohydrides, especially alkali metal cyanoborohydrides, for example sodium cyanoborohydride, in the presence of a suitable acid, preferably relatively weak acids, such as
- lower alkylcarboxylic acids, especially acetic acid, or a sulfonic acid, such as p-toluenesulfonic acid; in customary solvents, for example alcohols, such as MeOH or EtOH, or 20 ethers, for example cyclic ethers, such as THF, in the
- ethers, for example cyclic ethers, such as THF, in the presence or absence of water.

COCHECA DIECE

Scheme 2

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acid/amines 5 as shown in Scheme 2. Substituted amines are coupled with the mixed acid/amines 5 such as with a coupling reagent, for example EDC, to form the mixed amine/amide 6. Substituted carbonyl compounds, such as acid halides, anhydrides, carboxylic acids, esters, ketones, aldehydes and the like, are added to the mixed amine/amide 6 followed with reduction to give the substituted amide/substituted amine compounds 4.

Alternatively, compounds 4 can be prepared from mixed

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Scheme 3

Imino compounds 7 can be formed from the mixed amine/amides 6, such as by reacting with a substituted carbonyl compound.

10 Scheme 4

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FIGURATOR OFFICE

Substituted cyclic carboxamides can be prepared from
the corresponding imino analogs by the process outlined in
Scheme 4. Treatment of the imino compound 7 with a reducing
agent yields compound 4. Reagents which can be used to add
hydrogen to an imine double bond include borane in THF,
LiAlH4, NaBH4, sodium in EtOH and hydrogen in the presence
of a catalyst, and others.

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Scheme 5

Substituted carboxamides 4 can be prepared from the corresponding halo analogs 8 by the process outlined in Scheme 5. Substituted amino acids 9 are prepared from the corresponding chloro compounds 8 such as by reacting with an amine at a suitable temperature, such as about 80°C. The acid 9 is coupled with an amine, preferably in the presence of a coupling agent such as EDC, to form the corresponding amide 4.

The amination process can be carried out as an Ullmann type reaction using a copper catalyst, such as copper[0] or a copper[I] compound such as copper[I] coide, copper[I] bromide or copper[I] iodide in the presence of a suitable base (such as a metal carbonate, for example K2CO3) to neutralize the acid generated in the reaction. This reaction is reviewed in Houben-Weyl "Methoden der Organischen Chemie", Band 11/1, page 32 -33, 1958, in Organic Reactions, 14, page 19-24, 1965 and by J. Lindley

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(1984) in Tetrahedron, 40, page 1433-1456. The amount of catalyst is typically in the range of 1 to 20 mole percent. The reaction is carried out in an inert, aprotic solvent such as an ether (for example dimethoxyethane or dioxane) or an amide (for example dimethylformamide or N-methylpyrrolidone), under an inert atmosphere in the temperature range of $60\text{-}180^{\circ}\text{C}$.

An alternative amination process involves using a Group VIII element, where the metal core of the catalyst should be a zero-valent transition metal, such as palladium or nickel, which has the ability to undergo oxidative addition to the aryl-halogen bond. The zero valent state of the metal may be generated in situ from the M[II] state. The catalyst complexes may include chelating ligands, such as alkyl, aryl or heteroaryl derivatives of phosphines or biphosphines, imines or arsines. Preferred catalysts contain palladium or nickel. Examples of such catalysts include palladium[II]chloride, palladium[II]acetate, tetrakis(triphenyl-phosphine)palladium[0] and nickel[II]acetylacetonate. The metal catalyst is typically in the range of 0.1 to 10 mole percent. The chelating ligands may be either monodentate, as in the case for example of trialkyphosphines, such as tributylphosphine, triarylphosphines, such as tri-(ortho-tolyl)phosphine, and triheteroaryl phosphines, such as tri-2-furylphosphine; or they may be bidentate such as in the case of 2,2'-

bis(diphenylphosphino)ferrocene and 1-(N,N-dimethyl-amino)
1'-(dicyclohexylphosphino)biphenyl. The supporting ligand
may be complexed to the metal center in the form of a metal
complex prior to being added to the reaction mixture or may
be added to the reaction mixture as a separate compound. The
supporting ligand is typically present in the range 0.01 to

bis(diphenylphosphino) - 1,1'binaphthyl, 1,2-bis(diphenylphosphino)ethane, 1,1'-

20 mole percent. It is often necessary to add a suitable base to the reaction mixture, such as a trialkylamine (for example DIEA or 1,5-diazabicyclo[5,4,0]undec-5-ene), a Group I alkali metal alkoxide (for example potassium tert-butoxide) or carbonate (for example cesium carbonate) or potassium phosphate. The reaction is typically carried out in an inert aprotic solvent such as an ether (for example dimethoxyethane or dioxane) or an amide (for example, DMF or N-methylpyrrolidone), under an inert atmosphere in the temperature range of 60-180°C.

The amination is preferably carried out in an inert, aprotic, preferably anhydrous, solvent or solvent mixture, for example in a carboxylic acid amide, for example DMF or dimethylacetamide, a cyclic ether, for example THF or dioxane, or a nitrile, for example CH₂CN, or in a mixture thereof, at an appropriate temperature, for example in a temperature range of from about 40°C to about 180°C, and if necessary under an inert gas atmosphere, for example a nitrogen or argon atmosphere.

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Scheme 6

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Substituted carboxamides 4 can be prepared from the corresponding halo analogs 8 by the process outlined in Scheme 6. The chloro acid 8 is coupled with an amine, preferably in the presence of a coupling agent such as EDC, to form the corresponding chloro amide 10. Substituted amino-amides 4 are prepared from the corresponding chloro compounds 10 such as by reacting with an amine at a suitable temperature, such as about 80°C. The amination reaction can be run in the presence of an appropriate catalyst such as a palladium catalyst, in the presence of an aprotic base such as sodium t-butoxide or cesium carbonate, or a nickel catalyst, or a copper catalyst.

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Scheme 7

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Substituted carboxamides 4 can be prepared from the corresponding bromo/chloro analogs 11 by the process outlined in Scheme 7. The bromo/chloro acid 11 is coupled with an amine, preferably in the presence of a coupling agent such as EDC, to form the corresponding bromo substituted amide 12. Suzuki coupling with the bromo amide 12 and suitable boronic acids provides the substituted amide 10. Substituted amino-amides 4 are prepared from the corresponding chloro compounds 10 as described in Scheme 6.

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Scheme 8

Substituted thiophenes 16 can be prepared by the method of Scheme 8. The free amino group of a 3-amino-2-thiophenecarboxylic acid ester 13 can be protected such as by the addition of Boc_2O in a suitable solvent such as CH_2Cl_2 and DMAP. The ester is removed such as with base to form the free acid 14. The thiophene amide 15 is formed from the acid 14 such as by coupling with a substituted amine in the presence of DIEA, EDC and HOBt. The 2-protected-amino-

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thiophene amide 15 is deprotected, such as with 25% TFA/CH₂Cl₂. The free amine is alkylated such as with a substituted carboxaldehyde or similar active carbonyl compound, in the presence of a reducing agent NaCNBH₃ and the like, to form compounds 16.

Scheme 9

$$R^2$$
 N_{NH_2}
 $N_{$

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Substituted pyridines can be prepared such as by the method found in Scheme 9. 2-Aminonicotinic acid 17 is coupled with a substituted amine at a suitable temperature, nonprotic solvent such as CH₂Cl₂, such as with EDC and HOBt, to form the nicotinamide 18. The nicotinamide 18 is reductively alkylated such as with 4-pyridinecarboxaldehyde and NaBH(OAC)₃, to yield the 2-substituted amino-pyridyl carboxamides 19.

10

Schem 1

Substituted pyridines may be prepared by the method found in Scheme 10. 2-Chloro-nicotinic acid 20 is coupled with an amine 21 at a suitable temperature, such as a temperature over about 100°C to give the 2-substituted amino-nicotinic acid 22. The 2-substituted amino-nicotinic acid 20 is reacted with a substituted amine in the presence of a coupling reagent, such as BOP-Cl and base, such as TEA to form the 2-substituted amino-nicotinamide 19.

Alternatively, 2-chloro-nicotinoyl chloride (LG is Cl) is coupled first with R¹-NH₂ such as in the presence of base, e.g., NaHCO₃, in a suitable solvent, such as CH₂Cl₂, to form the amide 20A, then coupling with a pyridylmethylamine to yield the 2-substituted amino-nicotinamide 19.

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HODIOGE DESCRIPTION

Imino-substituted pyridines may be prepared by the method found in Scheme 11. (2-Amino-(4-pyridyl))-carboxamide 23 is reacted with 4-pyridine-carboxaldehyde, such as in the presence of p-toluenesulfonic acid monohydrate to yield the imino compound 24.

Scheme 12

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Substituted pyridines alternatively may be prepared by the method found in Scheme 12. The imino compound ${\bf 24}$ is reduced, such as with NaBH4, to form the substituted amine ${\bf 25}$.

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Scheme 13

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Substituted pyridines can be prepared by the process outlined in Scheme 13. A solution of sodium hypobromide is freshly prepared and added to 2-hydroxynicotinic acid 26 and heated, preferably at a temperature at about 50°C.

Additional sodium hypobromide may be needed to form the bromo compound 27. The 5-bromo-2-hydroxynicotinic acid 27 is reacted with thionyl chloride, preferably at a temperature >RT, more preferably at about 80°C to form the

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2-chloro-nicotinic acid analog 28. The acid is coupled with an amine, preferably in the presence of EDC, HOBT, and DIEA to form the corresponding substituted amide 29. Suzuki coupling with the brome amide and suitable boronic acids, provides the substituted nicotinamide 30. 2-Aminonicotinamides 31 are prepared from the corresponding chloro compounds 30 such as by reacting with substituted amines at a suitable temperature, such as about 80°C.

10 Scheme 14

Sulfonamides 32 can be prepared from amines 6 as shown in Scheme 14. Substituted sulfonyl compounds, such as sulfonyl halides, preferably chloro or bromo, sulfonic acids, an activated ester or reactive anhydride, or in the form of a cyclic amide, and the like, are added to the amine 6 to give the sulfonamide compounds 32.

The reaction is carried out in a suitable solvent, such as CH_2Cl_2 , at a temperature between about RT to about the reflux temperature of the solvent, in the presence of a suitable base, such as DIEA or DMAP.

The amino group of compounds 6 is preferably in free form, especially when the sulfonyl group reacting therewith is present in reactive form. The amino group may, however, itself be a derivative, for example by reaction with a phosphite, such as diethylchlorophosphite, 1,2-phenylene chlorophosphite, ethyldichlorophosphite, ethylene chlorophosphite or tetraethylpyrophosphite. A derivative of

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such a compound having an amino group also can be a carbamic acid halide or an isocvanate.

The condensation of activated sulfonic esters, reactive anhydrides or reactive cyclic amides with the corresponding amines is customarily carried out in the presence of an inorganic base, such as an alkaline metal hydrogen carbonate of carbonate, or especially an organic base, for example simple lower (alkyl)₃-amines, for example TEA or tributylamine, or one of the above-mentioned organic bases. If desired, a condensation agent is additionally used, for example as described for free carboxylic acids.

The condensation is preferably carried out in an inert, aprotic, preferably anhydrous, solvent or solvent mixture, for example in a carboxylic acid amide, for example formamide or DMF, a halogenated hydrocarbon, for example CH₂Cl₂, CCl₄ or chlorobenzene, a ketone, for example acetone, a cyclic ether, for example THF or dioxane, an ester, for example EtOAc, or a nitrile, for example CH₂CN, or in a mixture thereof, as appropriate at reduced or elevated temperature, for example in a temperature range of from about -40°C to about +100°C, preferably from about -10°C to about 70°C, and when arylsulfonyl esters are used, also at temperatures of from about 10-30°C, and if necessary under an inert gas atmosphere, for example a nitrogen or argon atmosphere.

Alcoholic solvents, for example EtOH, or aromatic solvents, for example benzene or toluene, may also be used. When alkali metal hydroxides are present as bases, acetone may also be added where appropriate.

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Scheme 15

Substituted pyridines can be prepared by the process outlined in Scheme 15. 2-Chloronicotinic acid 33 and substituted amine are coupled under conditions similar to that described in the previous schemes to give the amide 34. 6-Chloro-2-aminopyridines 35 are prepared from the amide 34, such as by reacting with substituted amines at a suitable temperature, such as above about 80°C, preferably above about 100°C, more preferably at about 130°C, neat. 6-Chloro-2-aminopyridines 35 are de-chlorinated such as by hydrogenation, for example by treatment with H₂ in the presence of Pd/C, to yield other compounds of the present invention 36.

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Scheme 16

1,2,3,6-Tetrahydro-pyridyl substituted anilines are prepared such as by the procedure described in Scheme 16 (where R^{x} is a substituent selected from those available for substituted R1). Nitrobenzenes 37 are brominated, such as with bromine in the presence of acid, H2SO4 for example, or with NBS to yield the 3-bromo derivative 38. Suzuki coupling of the bromo-derivative 38 and a substituted pyridylboronic acid, in an appropriate solvent such as toluene, such as at a temperature above RT, preferably above about 50°C, and more preferably at about 80°C, yields the pyridyl derivative 39. Alkylation of the nitrophenylpyridine 39, such as by treatment with iodomethane, preferably above about 50°C, and more preferably at about 80°C, yields the pyridinium compound 40, which upon reduction, such as by NaBH, yields the tetrahydyropyridine 41.

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Scheme 17

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6-Amino substituted pyridines are prepared such as by the procedure described in Scheme 17. Similar to the method of Scheme 13, chloropyridine 42 and is reacted with an amine, preferably above about 50°C, and more preferably at about 80°C, to yield the 6-aminopyridines 43.

Scheme 18

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A series of substituted anilines are prepared such as by the procedure described in Scheme 18. A nitrobenzyl bromide 44 is coupled with morpholine, such as at a temperature at about RT, to yield the heterocyclylmethyl nitrobenzene derivative. Reduction of the nitro compound, such as with iron powder, preferably above about 50°C, and more preferably at about 80°C, yields the heterocyclylmethyl substituted aniline 45.

Protected alkylamine substituted anilines can be prepared from the nitro free amines 46, such as with standard protecting agents and chemistry known in the art, such as BOC chemistry. Reduction of the protected nitro compound, such as with iron powder, preferably above about 50°C, and more preferably at about 80°C, yields the aniline 47.

Sulfonamide substituted anilines can be prepared from nitrobezenesulfonyl chlorides 48. Coupling of nitrobezenesulfonyl chlorides 48 with reactive heterocyclic compounds, such as substituted piperazines, piperidines, and the like, in a protic solvent such as EtOH, such as at a temperature about RT, yields the nitrobezenesulfonamides 48. Reduction of the nitro benzenesulfonamide, such as with iron powder, preferably above about 50°C, and more preferably at about 80°C, yields the aniline 49.

Scheme 19

A series of perhaloalkyl-substituted anilines ${\bf 52}$, where ${\bf R}^y$ represents perhaloalkyl radicals, are prepared such

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as by the procedure described in Scheme 19. 1-Nitro-4-(perfluoroethyl)benzene can be synthesized by the method described in the reference [John N. Freskos, Synthetic Communications, 18(9), 965-972 (1988)]. Alternatively, 1-Nitro-4-(perfluoroalkyl)benzene can be synthesized from the nitro compound, where X* is a leaving group, such as iodo, by the method described by W. A. Gregory, et al. [J. Med. Chem., 1990, 33, 2569-2578].

Reduction of the nitrobenzenes 51, such as with iron powder, at a temperature above about $50\,^{\circ}\text{C}$, and preferably at about $80\,^{\circ}\text{C}$, yields the aniline 52. Hydrogenation, such as with H_2 in the presence of catalyst, such as Pd/C, is also possible.

Scheme 20

Additional series of substituted anilines are prepared such as by the procedures described in Scheme 20 (where R^x is a substituent selected from those available for substituted R^1). 2-Alkoxy substituted anilines 55 are

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prepared from the corresponding phenol compounds 53 such as by the Mitsunobu reaction, including treatment with a N,N-dialkylethanolamine and PPh₃ and DEAD to give the corresponding nitro compound 54, followed by hydrogenation, such as with H, to give the aniline 55.

Alternatively, piperazinyl substituted anilines 58 can be prepared by the treatment of an aniline 56 with an N-substituted-bis(2-chloroethyl)amine, base, such as K_2CO_3 and NaI, at a temperature above about $50^{\circ}C$, preferably above about $100^{\circ}C$, and more preferably at about $170^{\circ}C$, to give the piperazinylbenzene compound 57. Nitration, such as with H_2SO_4 and HNO_3 , at a temperature above $0^{\circ}C$, and preferably at about RT, followed by hydrogenation, such as with H_2 atmosphere gives the substituted aniline 58.

Alternatively, piperazinyl substituted anilines **61** can be prepared by the treatment of a fluoro-nitro-substituted aryl compounds **59**. The fluoro-nitro-substituted aryl compounds **59** and 1-substituted piperazines are heated, preferably neat, at a temperature above about 50°C, and preferably at about 90°C, to yield the piperazinyl-nitroaryl compounds **60**. Hydrogenation, such as with H₂ atmosphere in the presence of a catalyst, such as 10% Pd/C, gives the substituted aniline **61**.

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Substituted indolines are prepared such as by the procedures described in Scheme 21. Substituted aminoindolines 64 are prepared from the nitroindoline 62 and a ketone in the presence of NaHE(OAc)₃ to form the 1-substituted indoline 63. The nitroindoline 63 is hydrogenated, such as with H_2 in the presence of a catalyst, such as Pd/C, to yield the amino-indoline 64.

Alternatively, substituted amino-indolines 67 are prepared from the nitroindoline 62. Nitroindoline 62, is reacted with an acid chloride to form an amide. Further treatment with a primary or secondary amine, preferably a secondary amine, such as in the presence of NaI, at a temperature above about $50\,^{\circ}\text{C}$, and preferably at about $70\,^{\circ}\text{C}$ yields the nitroindoline 65. The nitro compound 65 is hydrogenated, such as with H_2 in the presence of a catalyst, such as Pd/C, to yield the amino-indoline 66. The carbonyl is reduced, such as with BH₃-THF yields 1-aminoalkylindolines 67.

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Scheme 22

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Substituted indolines are prepared such as by the procedures described in Scheme 22. Substituted acetamides 69 are prepared from the acylation of halo-5-nitroanilines 68 (where LG is bromo or chloro, preferably chloro) with an acylating agent, such as acetyl chloride or acetic anhydride, under standard coupling chemistry, such as with DIEA, and DMAP, at a temperature of about RT, in a suitable solvent, such as CH₂Cl₂, DMF and/or DMAC. The N-(2-methylprop-2-enyl)acetamide 70 is prepared from the acetamide 69, such as by the treatment of base, such as NaH in anhydrous DMF and a 3-halo-2-methylpropene such as 3-bromo-2-methylpropene or 3-chloro-2-methylpropene, at a temperature between about 0°C and RT, and preferably at about RT; or with CsCO₃ at a temperature above RT, preferably above about 50°C and more preferably above about

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60°C. Cyclization of the N-(2-methylprop-2-enyl)acetamide 70, such as by the Heck-type reaction (treatment with Pd(OAc), in the presence of base, for example tetraethylammonium chloride, sodium formate, and NaOAc) at a temperature above about 50°C, and preferably at about 80°C. 5 yields the protected (3,3-dimethyl-2,3-dihydro-indol-1vl)ethanone 71. Deprotection, such as with strong acid such as AcOH on HCl at a temperature above about 50°C, and preferably at about 70-80°C, yields the 3,3-dimethyl-6-10 nitro-2,3-dihydro-indol-1-yl 72. Alternatively, the protected dihydro-6-nitro indoline 71 can be reduced, such as with Fe, or with 10% Pd/C in the presence of an excess of $\mathrm{NH_4CO_2H}$, or with $\mathrm{H_2}$ in the presence of a catalyst to form the protected dihydro-6-amino indoline 71a.

Scheme 23

Substituted anilines are prepared such as by the procedures described in Scheme 23. Nitrophenyl esters 74 are formed from the acid 73, such as by treatment with MeOH and acid. Alkylation of the ester 74, such as by treatment with base, followed by alkyl halide, yields the branched

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alkyl compounds 75. Reduction of the ester 75, such as with BH₃, yields the alcohol 76. The aldehyde 77 is prepared from the alcohol 76, such as by treatment with TPAP in the presence of N-methylmorpholine-N-oxide. Subsequent treatment with methoxymethyltriphenylphosphonium chloride and KHMDS yields 77. Coupling of the aldehyde 77 with morpholine, such as with NaBH(OAC)₃ yields the tertiary amine 78. Reduction of the nitro compound, such as with acid, for example AcOH, and zinc yields the aniline 79.

Scheme 24

Substituted aminomethyl compounds are prepared such as by the procedure described in Scheme 24. A piperidinemethanol 80 is reacted with formaldehyde and NaCNBH; Subsequently, base, such as sodium hydride, and a halo substituted cyclic nitrile gives the ether 81.

20 Hydrogenation of 81 under conditions described above, furnishes the aminomethyl compound 82.

Scheme 25

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Substituted aniline compounds are prepared such as by the procedure described in Scheme 25 (where $\ensuremath{R^{x}}$ is a substituent selected from those available for substituted \mathbb{R}^1 , preferably haloalkyl or alkyl). Alkynyl-aniline $\mathbf{84}$, prepared similar to that described in Scheme 46, is hydrogenated such as with H_2 in the presence of a catalyst, such as $Pd(OH)_2$, to yield the substituted alkyl 85.

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Scheme 26

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Substituted bromophenyl compounds are prepared such as by the procedure described in Scheme 26. Bromine is added to a optionally substituted nitrobenzene 86, silver(II)sulfate and acid, such as $\rm H_2SO_4$, to provide the bromo derivative 87.

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Scheme 27

Substituted anilines are prepared such as by the procedure described in Scheme 27 (where Rt and R are alkyl, or together with the nitrogen atom form a 4-6 membered heterocyclic ring). Acryloyl chloride 88 is reacted with an amine, preferably a secondary amine, such as at a temperature between about 0°C and about RT, to form the amide 89. A bromo-nitrobenzene 87 is reacted with the amide 89, such as in the presence of base, for example TEA, together with Pd(OAc)2 and Pd(PPh3)4, at a temperature above about 50°C, and preferably at about 120°C, such as in a sealed container, to form the substituted alkene 90. Hydrogenation of the alkene 90, such as with H2-in the presence of a catalyst, for example Pd/C catalyst yields the substituted aniline 91. Reduction of the amide 91, such as with LiAlH, at a temperature above about 50°C, and preferably at about 80°C yields the aniline 92.

Scheme 28

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Substituted indoles are prepared such as by the procedure described in Scheme 28. A nitroindole 93 is coupled with a halo compound, in the presence of base, for example $K_2\text{CO}_3$. Heating at a temperature above about $50\,^{\circ}\text{C}$, and preferably at about reflux yields the substituted-nitro-1H-indole 94. Hydrogenation similar to conditions described above yield the amino derivative 95.

Scheme 29

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Substituted pyrimidines are prepared such as by the procedure described in Scheme 29. 2-Methylthio-5-pyrimidyl acids 98 are prepared from the corresponding esters 96 similar to procedures described above. The amides 99 are formed from the acids 98 by coupling with the amine such as in the presence of HATU and base, TEA for example. The methylthio group can be removed, such as with Raney-Ni and heat, preferably at about reflux temperature, to form the pyrimidine 100.

Scheme 30

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Substituted aminomethyl compounds are prepared such as by the procedure described in Scheme 30 (where LG is a leaving group, such as Cl). Strong base, such as NaH is added to an alcohol and heated at about 50°C to form the sodium alkoxide, which is added to a halo compound, such as 2-chloro-4-cyanopyridine and heated at a temperature above about 50°C, and preferably at about 70°C to form the ether 102. Hydrogenation yields the aminomethyl derivative 103.

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Scheme 31

Substituted anilines are prepared such as by the procedure described in Scheme 31. Treatment with the haloalkyl alcohol 104 with an alcohol, such as in the presence of DEAD and PPh₃ yields the ether 105 or 106.

Scheme 32

$$\begin{array}{c|c}
 & LDA/CO_2 \\
\hline
 & N \\
\hline
 & F \\
\hline
 & 107 \\
\hline
 & 108 \\
\hline
 & 109 \\
\hline
\end{array}$$

Functionalized pyridines are prepared such as by the procedure described in Scheme 32. 2-Fluoropyridine 107 is treated with base, such as LDA at a temperature below about 0°C, and preferably at about -78°C, and quenched with a stream of dry CO₂ to form the nicotinic acid 108. Alternatively, solid CO₂ (dry ice) can be used, preferably dried with N₂ prior to use. The acid 108 is converted to the acid halide 109, such as by treatment with thionyl chloride and heating at a temperature above about 50°C, and preferably at about reflux.

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Scheme 33

Chloro-substituted pyridines 110 are prepared such as by the procedure described in Scheme 33. 2-Chloronicotinic acid is activated with ethyl chloroformate, in the presence of base, such as TEA, at a temperature of about RT. Reaction with an amine produces amide 110. Alternatively, the amine can be coupled with the acid chloride 111, such as with polymer-supported DIPEA, to form amide 110. Excess acid chloride is removed by treating the reaction mixture with polymer-supported trisamine resin.

Scheme 34

Amino-substituted indoles 110 are prepared such as by the procedure described in Scheme 34. Nitroindoline 112 is reacted with N-methyl-4-piperidone in the presence of NaOMe at a temperature above about 50°C, and preferably at about reflux, to form the 3-substituted indole 113. Hydrogenation as previously discussed yields the amino indole 114.

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$$O_2N \longrightarrow \stackrel{H}{\stackrel{N}{\longrightarrow}} \stackrel{N}{\stackrel{N}{\longrightarrow}} \stackrel{NaH, R^*I}{\longrightarrow} O_2N \longrightarrow \stackrel{R^*}{\stackrel{N}{\longrightarrow}} \stackrel{H_2}{\longrightarrow} \stackrel{H_2N}{\longrightarrow} \stackrel{R^*}{\longrightarrow} \stackrel{R^*}{\longrightarrow} \stackrel{N}{\longrightarrow} \stackrel$$

Alkylated indazoles can be prepared by the process outlined in Scheme 35. To a solution of 6-nitroindazole 115 in a solvent such as THF is added strong base, such as NaH at a temperature below RT, preferably at about 0°C. Alkylhalides, such as where R" is methyl, are added and reacted at a temperature about RT to give 1-alkyl-6-nitro-1H-indazole 116. The nitro indazole 116 is hydrogenated, such as with an H₂ atmosphere in the presence of a catalyst, such as Pd/C to give the 1-substituted-6-amino-1H-indazole

Scheme 36

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Brominated indazoles can be prepared by the process outlined in Scheme 36. NBS is slowly added to an acidic solution, such as a mixture of TFA:H₂SO₄ (5:1) and tert-butyl-4-nitrobenzene 118 at a temperature of about RT to yield the brominated compound 119.

Schem 37

Substituted anilines can be prepared by the process 5 outlined in Scheme 38. A mixture of 1-(substituted)-2bromo-4-nitrobenzene 120 (where Rx is a substituent selected from those available for substituted R1) and Nmethylpiperazine is heated, such as with or without solvent, preferably without solvent, at a temperature above RT, 10 preferably at a temperature above about 100°C, and more preferably at a temperature at about 130°C to give the 1-[5-(substituted) -2-nitrophenyl]-4-methylpiperazine 121. The nitro compound 121 is hydrogenated, such as with an H2 atmosphere in the presence of a catalyst, such as Pd/C to 15 furnish 4-(substituted)-2-(4-methylpiperazinyl)phenylamine 122.

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Scheme 38

Tricyclic heterocycles can be prepared by the process outlined in Scheme 38. 7-Nitro-2,3,4-trihydroisoquinolin-1-one 123 is heated in FOCl₃ at a temperature above RT, preferably at a temperature sufficient for reflux, to form the 1-chloro-7-nitro-3,4-dihydroisoquinoline 124. The 1-chloro-7-nitro-3,4-dihydroisoquinoline 124 is dissolved in a solvent, such as THF, and H₂NNH₂ is added. The reaction is evaporated to a residue, then heated with HC(OEt)₃ at a temperature above RT, preferably at a temperature above about 75°C, and more preferably at a temperature at about 115°C to give the nitro-substituted tricyclic. Hydrogenation, such as with an H₂ atmosphere in the presence of a catalyst, such as Fd/C, gives 2-amino-5,6,7-trihydro-1,2,4-triazolo(3,4-a)isoquinoline 125.

Scheme 39

128

5 Indazolvl ethers can be prepared by the process outlined in Scheme 39. 6-Nitro-1H-2-hvdroindazol-3-one 126 is protected such as with Boc2O and DMAP in CH2Cl2 at a temperature of about RT, to give the protected 6-nitro-2hydroindazol-3-one. The protected 6-nitro-2-hydroindazol-3-10 one is reacted with an alcohol (where Rx is an appropriate substituent selected from the possible substituents on R) and Ph₃P in a solvent, such as THF, and DEAD, at a temperature of about RT, to give the protected 6nitro(indazol-3-vl) ether. The nitro intermediate is hydrogenated, such as with an ${\rm H}_2$ atmosphere in the presence 15 of a catalyst, such as Pd/C, to give the protected 6amino(indazol-3-yl) ether 127. The amine 127 is coupled and 2-chloronicotinic acid in a solvent, such as an alcohol, preferably pentanol, at a temperature above RT, preferably 20 at a temperature above about 75°C, and more preferably at a

temperature at about $130\,^{\circ}\text{C}$ to give the coupled and deprotected compound 128.

Indolinyl substituted carboxamides can be prepared from the corresponding nitro indoline 129 by the process outlined in Scheme 40. For example, 3,3-dimethyl-6-nitroindoline 129 is alkylated, such as with N-protected-4-formylpiperidine in the presence of NaHB(OAc), and acid, such as glacial AcOH, and solvent, such as dichloromethane, at a temperature of about RT, to afford the alkylated indane 130. Hydrogenation of the alkylated indane 130, such as

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with an H₂ atmosphere in the presence of a catalyst, such as Pd/C, in the presence of a solvent, such as an alcohol, preferably MeOH, to give the amino intermediate 131. Alternatively, other hydrogenation methods can be used, such as Fe powder with NH₄Cl. Coupling of the amine 131, such as with 2-chloronicotinic acid and DIEA, HOBt and EDC, in a solvent such as CH₂Cl₂ at a temperature of about RT provides the protected carboxamide 132, which upon deprotection and alkylation yields other compounds of the invention, 133 and 134, respectively. Alternatively, amine 131 is reacted with 2-fluoronicotinoyl chloride to form a 2-fluoronicotinamide, which can be alkylated, such as in Scheme 10.

Scheme 41

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Substituted anilines can be prepared by the process outlined in Scheme 41. 1-Methyl-4-piperidinone 135 is added

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such as THF, at a temperature below RT, preferably lower than about -50°C, more preferably at about -78°C. TfoNPh is reacted with the enolate at a temperature of about RT, to give 1-methyl-4-(1,2,5,6-tetrahydro)pyridyl-(trifluoromethyl)sulfonate. A mixture of the triflate intermediate, bis(pinacolato)diboron, potassium acetate, PdClodppf, and dppf in a solvent such as dioxane is heated at a temperature above RT, preferably at a temperature above about 50°C, and more preferably at a temperature at about 10 80°C to give 4.4.5.5-tetramethyl-2-(1-methyl(4-1.2.5.6tetrahydropyridyl))-1,3,2-dioxaborolane 136. The substituted aniline 137 is formed from the 1,3,2dioxaborolane 136 such as with treatment with an amine in 15 the presence of 1,1'-bis(diphenyphosphino)ferrocenepalladium dichloride and base, such as K2CO2, in a solvent such as DMF at a temperature above RT, preferably at a temperature above about 50°C, and more preferably at a temperature at about 80°C.

to a solution of strong base such as LiHMDS. in a solvent

Scheme 42

138 KOH,
$$\Delta$$
 alkylation 140

nitration v

hydrogenation 02N

141

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Substituted anilines can be prepared by the process outlined in Scheme 42. 4-Cvano-4-phenylpiperidine hydrochloride 138 is treated with base, such as KOH, at a temperature above RT, preferably at a temperature above about 100°C, and more preferably at a temperature at about 160°C, to provide the phenyl piperidine 139. Alkylation of the phenyl piperidine 139, such as with formaldehyde and NaCNBH, in a solvent such as CH,CN, with sufficient acid to maintain the reaction pH near 7, to provide the alkylated piperidine 140. Nitration of the phenylpiperidine 140, such as with H2SO, and fuming HNO2 at a temperature below RT, and preferably at about 0°C, gives the nitro intermediate 141. Hydrogenation of the nitro intermediate 141, such as with an H₂ atmosphere in the presence of a catalyst, such as Pd/C, in the presence of a solvent, such as an alcohol, preferably MeOH, to give the amino intermediate 142.

Scheme 43

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Substituted amides can be prepared by the process outlined in Scheme 43. 3-Nitrocinnamic acid 143 is coupled with 1-methylpiperazine in the presence of EDC and a solvent such as CH2Cl2, at a temperature of about RT gives the carboxamide 144.

Scheme 44

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Substituted benzylamines can be prepared by the process outlined in Scheme 44. A substituted bromobenzylamine 145 where R^{1a} is a substituent described for R¹ is protected such as with Boc₂O in the presence of base, such as TEA in an appropriate solvent such as CH₂Cl₂. The protected bromobenzylamine 146 is alkylated, such as with 1-dimethylamino-2-propyne in the presence of catalyst, such as PdCl₂(PPh₃)₂, and CuI, in the presence of base, such as TEA, at a temperature above RT, preferably at a temperature above about 50°C, and more preferably at a

- 15 temperature above about 50°C, and more preferably at a temperature at about 100°C, such as in a sealed tube, to form the propynylbenzylamine 147. The propynylbenzylamine is hydrogenated such as with $\rm H_2$ in the presence of Pd(OH) $_2$ and MeOH to provide the propylbenzylamine 148.
- 20 Deprotection, such as with strong acid, such as TFA, for removal of a Boc protecting group, yields the propylbenzylamine 149.

Scheme 45

Substituted benzylamines can be prepared by the process outlined in Scheme 45. The protected bromobenzylamine 146 is alkylated, such as with propargyl alcohol in the presence of catalyst, such as PdCl₂(PPh₃), and CuI, in the presence of base, such as TEA, at a temperature above RT, preferably at a temperature above about 50°C, and more preferably at a temperature at about 100°C, such as in a sealed tube, to form the protected hydroxypropynylbenzylamine 150. The protected hydroxypropynylbenzylamine is treated with N-methylmorpholine oxide in the presence of a catalyst, such as tetrapropylammonium perruthenate, to form the aldehyde

addition of morpholine and NaBH(OAc)₃ provides the morpholinyl derivative. Deprotection, such as with strong acid, such as TFA, for removal of a Boc protecting group, yields the propylbenzylamine 151.

intermediate. Reductive amination, such as with the

Scheme 46

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Substituted aminomethyl compounds are prepared such as by the procedure described in Scheme 46. A halo compound 152, is reacted with an alkyne in the presence of PdCl₂(PPh₃)₂ and CuI, with base is heated at a temperature above about 50°C, and preferably at about 100°C, such as in a sealed container, to provide the substituted alkyne 153.

Scheme 47

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Substituted heterocycles may be prepared by the method found in Scheme 47. Chloro-heterocycles 154 (where LG is OH) is coupled with an amine 155 at a suitable temperature, such as a temperature over about 100°C to give the 2-substituted amino-nicotinic acid 156. The 2-substituted amino-nicotinic acid 156 is reacted with a substituted amine in the presence of a coupling reagent, such as BOP-Cl and base, such as TEA to form the 2-substituted amino-nicotinamide 157.

Alternatively, 2-chloro-nicotinoyl chloride 154 (where LG is Cl) is coupled first with $R^2\text{-NH}_2$, such as in the presence of base, e.g., NaHCO₃, in a suitable solvent, such

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as IpOH or CH₂Cl₂, to form the amide 158, then coupled with an amine 155 to yield the 2-substituted amino-nicotinamide 157. Where A is a pi-electron rich heterocycle, the addition of KF, such as 40% KF on alumina in IpOH, at a temperature over about 100°C, preferably about 160°C, can be used in the formation of 157 from 158.

Scheme 48

2,3,4,4a,9,9a-hexahydro-1H-3-aza-fluoren-6-ylamine may be prepared by the method found in Scheme 48.

Nitrobenzylpyridines 159 are alkylated, such as with MeI, in 15 the presence of TBAI and base to form the pyridinium compound 160. The pyridinium compounds 160 are halogenated, such as brominated with NBS, to form the brominated pyridinium compounds 161 which are reduced such as with

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NaBH₄ to form the tetrahydro-pyridines 162. Palladium catalyzed intramolecular Heck coupling followed by hydrogenation forms the hexahydro-fluorenes 164.

The starting compounds defined in Schemes 1-48 may also be present with functional groups in protected form if necessary and/or in the form of salts, provided a saltforming group is present and the reaction in salt form is possible. If so desired, one compound of formulas I-XII can be converted into another compound of formulas I-XII or a Noxide thereof; a compound of formulas I-XII can be converted into a salt; a salt of a compound of formulas I-XII can be converted into the free compound or another salt; and/or a mixture of isomeric compounds of formulas I-XII can be separated into the individual isomers.

N-Oxides can be obtained in a known matter by reacting a compound of formulas I-XII with hydrogen peroxide or a peracid, e.g. 3-chloroperoxy-benzoic acid, in an inert solvent, e.g. dichloromethane, at a temperature between about -10-35°C, such as about 0°C - RT.

If one or more other functional groups, for example carboxy, hydroxy, amino, or mercapto, are or need to be protected in a compound of formulas I-XII or in the synthesis of a compound of formulas I-XII, because they should not take part in the reaction, these are such groups as are usually used in the synthesis of peptide compounds, and also of cephalosporins and penicillins, as well as nucleic acid derivatives and sugars.

The protecting groups may already be present in precursors and should protect the functional groups concerned against unwanted secondary reactions, such as acylations, etherifications, esterifications, oxidations, solvolysis, and similar reactions. It is a characteristic of protecting groups that they lend themselves readily, i.e.

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without undesired secondary reactions, to removal, typically by solvolysis, reduction, photolysis or also by enzyme activity, for example under conditions analogous to physiological conditions, and that they are not present in the end-products. The specialist knows, or can easily establish, which protecting groups are suitable with the reactions mentioned above and hereinafter.

The protection of such functional groups by such protecting groups, the protecting groups themselves, and their removal reactions are described for example in standard reference works, such as J. F. W. McOmie, "Protective Groups in Organic Chemistry", Plenum Press, London and New York 1973, in T. W. Greene, "Protective Groups in Organic Synthesis", Wiley, New York 1981, in "The Peptides": Volume 3 (editors: E. Gross and J. Meienhofer). Academic Press, London and New York 1981, in "Methoden der organischen Chemie" (Methods of organic chemistry), Houben Weyl, 4th edition, Volume 15/1, Georg Thieme Verlag, Stuttgart 1974, in H.-D. Jakubke and H. Jescheit, "Aminosauren, Peptide, Proteine" (Amino acids, peptides, proteins), Verlag Chemie, Weinheim, Deerfield Beach, and Basel 1982, and in Jochen Lehmann, "Chemie der Kohlenhydrate: Monosaccharide und Derivate" (Chemistry of

In the additional process steps, carried out as desired, functional groups of the starting compounds which should not take part in the reaction may be present in unprotected form or may be protected for example by one or more of the protecting groups mentioned above under "protecting groups". The protecting groups are then wholly or partly removed according to one of the methods described there.

carbohydrates: monosaccharides and derivatives), Georg

Thieme Verlag, Stuttgart 1974.

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Salts of a compound of formulas I-XII with a saltforming group may be prepared in a manner known per se. Acid
addition salts of compounds of formulas I-XII may thus be
obtained by treatment with an acid or with a suitable anion
exchange reagent. A salt with two acid molecules (for
example a dihalogenide of a compound of formulas I-XII) may
also be converted into a salt with one acid molecule per
compound (for example a monohalogenide); this may be done by
heating to a melt, or for example by heating as a solid
under a high vacuum at elevated temperature, for example
from about 130°C to about 170°C, one molecule of the acid
being expelled per molecule of a compound of formulas I-XII.

Salts can usually be converted to free compounds, e.g. by treating with suitable basic agents, for example with alkali metal carbonates, alkali metal hydrogen carbonates, or alkali metal hydroxides, typically potassium carbonate or sodium hydroxide.

A compound of formulas I-XII, wherein Z is oxygen, can be converted into the respective compound wherein Z is sulfur, for example, by using an appropriate sulfur compound, e. g. using reaction with Lawesson's reagent (2,4-bis-(4-methoxyphenyl)-1,3-dithia-2,4-diphosphetane-2,4-disulfide) in a halogenated hydrocarbon, such as CH₂Cl₂, or an aprotic solvent, such as toluene or xylene, at temperatures from about 30°C to reflux.

All process steps described here can be carried out under known reaction conditions, preferably under those specifically mentioned, in the absence of or usually in the presence of solvents or diluents, preferably such as are inert to the reagents used and able to dissolve these, in the absence or presence of catalysts, condensing agents or neutralizing agents, for example ion exchangers, typically cation exchangers, for example in the H+ form, depending on the type of reaction and/or reactants at reduced, normal, or

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elevated temperature, for example in the range from about -100°C to about 190°C, preferably from about -80°C to about 150°C, for example at about -80 to about 60°C, at RT, at about -20 to about 40°C or at the boiling point of the solvent used, under atmospheric pressure or in a closed vessel, where appropriate under pressure, and/or in an inert atmosphere, for example under argon or nitrogen.

Salts may be present in all starting compounds and transients, if these contain salt-forming groups. Salts may also be present during the reaction of such compounds, provided the reaction is not thereby disturbed.

In certain cases, typically in hydrogenation processes, it is possible to achieve stereoselective reactions, allowing for example easier recovery of individual isomers.

The solvents from which those can be selected which are suitable for the reaction in question include for example water, esters, typically lower alkyl-lower alkanoates, e.g., ethyl acetate, ethers, typically aliphatic ethers, e.g., diethylether, or cyclic ethers, e.g., THF, liquid aromatic hydrocarbons, typically benzene or toluene, alcohols, typically MeOH, EtOH or 1-propanol, IPOH, nitriles, typically CH2CN, halogenated hydrocarbons, typically CH2Cl2, acid amides, typically DMF, bases, typically heterocyclic nitrogen bases, e.g. pyridine, carboxylic acids, typically lower alkanecarboxylic acids, e.g., AcOH, carboxylic acid anhydrides, typically lower alkane acid anhydrides, e.g., acetic anhydride, cyclic, linear, or branched hydrocarbons, typically cyclohexane, hexane, or isopentane, or mixtures of these solvents, e.g., aqueous solutions, unless otherwise stated in the description of the process. Such solvent mixtures may also be used in processing, for example in chromatography.

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The invention relates also to those forms of the process in which one starts from a compound obtainable at any stage as a transient and carries out the missing steps, or breaks off the process at any stage, or forms a starting material under the reaction conditions, or uses said starting material in the form of a reactive derivative or salt, or produces a compound obtainable by means of the process according to the invention and processes the said compound in situ. In the preferred embodiment, one starts from those starting materials which lead to the compounds described above as preferred.

The compounds of formulas I-XII, including their salts, are also obtainable in the form of hydrates, or their crystals can include for example the solvent used for crystallization (present as solvates).

New starting materials and/or intermediates, as well as processes for the preparation thereof, are likewise the subject of this invention. In the preferred embodiment, such starting materials are used and reaction conditions so selected as to enable the preferred compounds to be obtained.

Starting materials of the invention, are known, are commercially available, or can be synthesized in analogy to or according to methods that are known in the art.

For example, amine 1 can be prepared by reduction of the corresponding nitro. The reduction preferably takes place in the presence of a suitable reducing agent, such as tin(II) chloride or hydrogen in the presence of an appropriate catalyst, such as Raney nickel (then preferably the hydrogen is used under pressure, e.g. between 2 and 20 bar) or PtO₂, in an appropriate solvent, e.g. an alcohol, such as MeOH. The reaction temperature is preferably between about 0°C and about 80°C, especially about 15°C to about 30°C.

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It would also be possible to reduce the nitro compound after forming the amide compound under reaction conditions analogous to those for the reduction of nitro compounds described above. This would eliminate the need to protect the free amino group as described in Scheme 1.

In the preparation of starting materials, existing functional groups which do not participate in the reaction should, if necessary, be protected. Preferred protecting groups, their introduction and their removal are described above or in the examples.

All remaining starting materials are known, capable of being prepared according to known processes, or commercially obtainable; in particular, they can be prepared using processes as described in the examples.

Compounds of the present invention can possess, in

general, one or more asymmetric carbon atoms and are thus capable of existing in the form of optical isomers as well as in the form of racemic or non-racemic mixtures thereof. The optical isomers can be obtained by resolution of the racemic mixtures according to conventional processes, e.g., by formation of diastereoisomeric salts, by treatment with an optically active acid or base. Examples of appropriate acids are tartaric, diacetyltartaric, dibenzoyltartaric, ditoluoyltartaric, and camphorsulfonic acid and then separation of the mixture of diastereoisomers by crystallization followed by liberation of the optically active bases from these salts. A different process for separation of optical isomers involves the use of a chiral chromatography column optimally chosen to maximize the separation of the enantiomers. Still another available method involves synthesis of covalent diastereoisomeric molecules by reacting compounds of the invention with an optically pure acid in an activated form or an optically pure isocyanate. The synthesized diastereoisomers can be

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separated by conventional means such as chromatography, distillation, crystallization or sublimation, and then hydrolyzed to deliver the enantiomerically pure compound. The optically active compounds of the invention can likewise be obtained by using optically active starting materials. These isomers may be in the form of a free acid. a free base, an ester or a salt.

The compounds of this invention may contain one or more asymmetric centers and thus occur as racemates and racemic mixtures, scalemic mixtures, single enantiomers, individual diastereomers and diastereomeric mixtures. All such isomeric forms of these compounds are expressly included in the present invention.

The compounds of this invention may also be

15 represented in multiple tautomeric forms, for example, as
illustrated below:

20 The invention expressly includes all tautomeric forms of the compounds described herein.

The compounds may also occur in cis- or trans- or Eor Z- double bond isomeric forms. All such isomeric forms of such compounds are expressly included in the present invention. All crystal forms of the compounds described

25 invention. All crystal forms of the compounds described herein are expressly included in the present invention.

Substituents on ring moieties (e.g., phenyl, thienyl, etc.) may be attached to specific atoms, whereby they are intended to be fixed to that atom, or they may be drawn unattached to a specific atom, whereby they are intended to

be attached at any available atom that is not already substituted by an atom other than H (hydrogen).

The compounds of this invention may contain heterocyclic ring systems attached to another ring system. Such heterocyclic ring systems may be attached through a

carbon atom or a heteroatom in the ring system.

Alternatively, a compound of any of the formulas delineated herein may be synthesized according to any of the processes delineated herein. In the processes delineated

10 herein, the steps may be performed in an alternate order and may be preceded, or followed, by additional protection/deprotection steps as necessary. The processes may further comprise use of appropriate reaction conditions, including inert solvents, additional reagents, such as bases

15 (e.g., LDA, DIEA, pyridine, K₂CO₃, and the like), catalysts, and salt forms of the above. The intermediates may be isolated or carried on in situ, with or without purification. Purification methods are known in the art and include, for example, crystallization, chromatography

20 (liquid and gas phase, simulated moving bed ("SMB")), extraction, distillation, trituration, reverse phase HPLC and the like. Reactions conditions such as temperature, duration, pressure, and atmosphere (inert gas, ambient) are known in the art and may be adjusted as appropriate for the 25 reaction.

As can be appreciated by the skilled artisan, the above synthetic schemes are not intended to comprise a comprehensive list of all means by which the compounds described and claimed in this application may be synthesized. Further methods will be evident to those of ordinary skill in the art. Additionally, the various synthetic steps described above may be performed in an alternate sequence or order to give the desired compounds. Synthetic chemistry transformations and protecting group

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methodologies (protection and deprotection) useful in synthesizing the inhibitor compounds described herein are known in the art and include, for example, those such as described in R. Larock, Comprehensive Organic

- Transformations, VCH Publishers (1989); T.W. Greene and P.G.M. Wuts, Protective Groups in Organic Synthesis, 3rd. Ed., John Wiley and Sons (1999); L. Fieser and M. Fieser, Fieser and Fieser's Reagents for Organic Synthesis, John Wiley and Sons (1994); A. Katritzky and A. Pozharski,
- Handbook of Heterocyclic Chemistry, 2nd Ed. (2001); M. Bodanszky, A. Bodanszky: The practice of Peptide Synthesis Springer-Verlag, Berlin Heidelberg 1984; J. Seyden-Penne: Reductions by the Alumino- and Borohydrides in Organic Synthesis, 2nd Ed., Wiley-VCH, 1997; and L. Paquette, ed.,
 Encyclopedia of Reagents for Organic Synthesis, John Wiley
 - 5 Encyclopedia of Reagents for Organic Synthesis, John Wiley and Sons (1995).

The compounds of this invention may be modified by appending appropriate functionalities to enhance selective biological properties. Such modifications are known in the art and include those which increase biological penetration into a given biological compartment (e.g., blood, lymphatic system, central nervous system), increase oral availability, increase solubility to allow administration by injection, alter metabolism and alter rate of excretion.

The following examples contain detailed descriptions of the methods of preparation of compounds of Formulas I-XII. These detailed descriptions fall within the scope, and serve to exemplify, the above described General Synthetic Procedures which form part of the invention. These detailed descriptions are presented for illustrative purposes only and are not intended as a restriction on the scope of the invention.

Unless otherwise noted, all materials were obtained from commercial suppliers and used without further

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purification. Anhydrous solvents such as DMF, THF, CH_2Cl_2 and toluene were obtained from the Aldrich Chemical Company. All reactions involving air- or moisture-sensitive compounds were performed under a nitrogen atmosphere. Flash

- 5 chromatography was performed using Aldrich Chemical Company silica gel (200-400 mesh, 60A) or Biotage pre-packed column. Thin-layer chromatography (TLC) was performed with Analtech gel TLC plates (250 μ). Preparative TLC was performed with Analtech silica gel plates (1000-2000 μ). Preparative HPLC
- 10 was conducted on Beckman or Waters HPLC system with 0.1% TFA/H₂O and 0.1% TFA/CH₃CN as mobile phase. The flow rate was at 20 ml/min. and gradient method was used. ¹H NMR spectra were determined with super conducting FT NMR spectrometers operating at 400 MHz or a Varian 300 MHz
- 15 instrument. Chemical shifts are expressed in ppm downfield from internal standard tetramethylsilane. All compounds showed NMR spectra consistent with their assigned structures. Mass spectra (MS) were determined on a Perkin Elmer - SCIEX API 165 electrospray mass spectrometer
- 20 (positive and, or negative) or an HP 1100 MSD LC-MS with eletrospray ionization and quadrupole detection. All parts are by weight and temperatures are in Degrees centigrade unless otherwise indicated.

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The following abbreviations are used:

	AIBN -	2,2'-azobisisobutyronitrile
	Ar -	argon
5	AgSO ₄ -	silver sulfate
	ATP -	adenosine triphosphate
	BH ₃ -	borane
	Boc -	tert-butyloxycarbonyl
	Boc ₂ O -	Boc anhydride
10	BOP-C1 -	bis(2-oxo-3-oxazolidinyl)phosphinic
		chloride
	Br ₂ -	bromine
	BSA -	bovine serum albumin
	t-BuOH -	tert-butanol
15	CAN -	ammonium cerium(IV) nitrate
	CH3CN, AcCN -	acetonitrile
	CH ₂ Cl ₂ -	dichloromethane
	CH ₃ I, MeI -	iodomethane, methyl iodide
	CCl ₄ -	carbon tetrachloride
20	CCl ₃ -	chloroform
	CO ₂ -	carbon dioxide
	Cs ₂ CO ₃ -	cesium carbonate
	DIEA -	diisopropylethylamine
	CuI -	copper iodide
25	DCE -	1,2-dichloroethane
	DEAD -	diethyl azodicarboxylate
	DIEA -	diisopropylethylamine
	dppf -	1,1-diphenylphosphinoferrocene
	DMAP -	4-(dimethylamino)pyridine
30	DMAC -	N,N-dimethylacetamide
	DMF -	dimethylformamide
	DMSO -	dimethylsulfoxide
	DTT -	dithiothreitol

EDC, EDAC- 1-(3-dimethylaminopropyl)-3-

LiCl -

MeOH -

MgCl₂ -

LiHMDS -

		ethylcarbodiimide hydrochloride
	EGTA -	ethylene glycol-bis(β -aminoethyl ether)-
		N,N,N',N'-tetraacetic acid
	EtOAc -	ethyl acetate
5	EtOH -	ethanol
	Et ₂ O -	diethyl ether
	Fe -	iron
	g -	gram
	h -	hour
10	HATU -	O-(7-azabenzotriazol-1-yl)-N,N,N',N'-
		${\tt tetramethyluronium\ hexafluorophosphate}$
	H ₂ -	hydrogen
	H ₂ O -	water
	HCl -	hydrochloric acid
15	H ₂ SO ₄ -	sulfuric acid
	H ₂ NNH ₂ -	hydrazine
	HC(OEt) ₃ -	triethylorthoformate
	HCHO, H ₂ CO -	formaldehyde
	HCO₂Na -	sodium formate
20	HOAc, AcOH -	acetic acid
	HOAt -	1-hydroxy-7-azabenzotriazole
	HOBt -	hydroxybenzotriazole
	IpOH -	isopropanol
	K ₂ CO ₃ -	potassium carbonate
25	KHMDS -	potassium hexamethylsilazane
	KNO ₃ -	potassium nitrate
	KOAc -	potassium acetate
	КОН -	potassium hydroxide
	LAH, LiAlH4 -	lithium aluminum hydride
30	LDA -	lithium diisopropylamide

lithium chloride

magnesium chloride

methanol

lithium hexamethyldisilazide

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PtO₂ -

RT -

Maso, magnesium sulfate mar milligram m1 milliliter MnCl₂ manganese chloride NBS -N-bromosuccinimide NMO -4-methylmorpholine, N-oxide NMP -N-methylpyrrolidone Na₂SO₄ sodium sulfate Na₂S₂O₅ sodium metabisulfite 10 NaHCO₃ sodium bicarbonate Na₂CO₃ sodium carbonate NaCl sodium chloride NaH sodium hydride NaI sodium iodide 15 NaOH sodium hydroxide NaOMe sodium methoxide sodium cyanoborohydride NaCNBH3 -NaBH4 sodium borohydride NaNO2 sodium nitrate 20 NaBH (OAc) 3 sodium triacetoxyborohydride NH4Cl ammonium chloride N₂ nitrogen Pd/C palladium on carbon PdCl2(PPh3)2 palladium chloride bis(triphenylphosphine) 25 PdCl2(dppf) -1,1-bis(diphenvlphosphino)ferrocene palladium chloride Pd(PPh3)4 palladium tetrakis triphenylphosphine Pd(OH), palladium hydroxide Pd(OAc) 2 palladium acetate 30 PMB para methoxybenzyl POCl₁ phosphorus oxychloride PPh₃ triphenylphosphine

platinum oxide

room temperature

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SiO₂ - silica

SOCl₂ - thionyl chloride

TBAI - tetrabutylammonium iodide

TEA - triethylamine

5 Tf₂NPh - N-phenyltrifluoromethanesulfonimide

TFA - trifluoroacetic acid

THF - tetrahydrofuran

TPAP - tetrapropylammoniumperruthenate

Tris-HCl - Tris(hydroxymethyl)aminomethane

10 hydrochloride salt

Zn - zinc

Preparation I - 3-nitro-5-trifluoromethyl-phenol

1-Methoxy-3-nitro-5-trifluoromethyl-benzene (10g, Aldrich)

15 and pyridine-HCl (41.8g, Aldrich) were mixed together and
heated neat at 210°C in an open flask. After 2.5 h the
mixture was cooled to RT and partitioned between 1N HCl and
EtOAc. The EtOAc fraction was washed with 1N HCl (4x), brine
(1x), dried with Na₂SO₄, filtered and concentrated in vacuo

20 to form 3-nitro-5-trifluoromethyl-phenol as an off-white solid.

Preparation II - 1-Boc-4-(3-nitro-5-trifluoromethylphenoxy)-piperidine

25 3-Nitro-5-trifluoromethyl-phenol (8.81g) was dissolved in THF (76 ml). 1-Boc-4-hydroxy-piperidine (8.81 g, Aldrich) and Ph₃P (11.15 g) were added and the solution was cooled to -20°C. A solution of DEAD (6.8 ml, Aldrich) in THF (36 ml) was added dropwise, maintaining the temperature between -20 and -10°C. The reaction was warmed to RT and stirred overnight. The reaction was concentrated in vacuo and triturated with hexane. The yellow solid was removed by filtration and washed with Et₂O (25 ml), and hexane. The

white filtrate was washed with 1N NaOH (2x), brine (1x) and

the hexane layer was dried over Na_2SO_4 , filtered and concentrated in vacuo. The crude material was purified with flash chromatography (SiO₂, 5-10% EtOAc/hexane) to obtain 1-Boc-4-(3-nitro-5-trifluoromethyl-phenoxy)-piperidine.

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The following compounds were prepared similarly to the procedure outlined above:

- a) (S)-1-Boc-[2-(5-nitro-2-trifluoromethylphenoxymethyl]-10 pvrrolidine
 - b) (R)-1-Boc-[2-(5-nitro-2-trifluoromethylphenoxymethyl]pyrrolidine.
 - c) (R) 1-Boc-2-(3-Nitro-5-trifluoromethyl-phenoxymethyl)pyrrolidine
- 15 d) 4-(2-tert-Butyl-5-nitro-phenoxymethyl)-1-methylpiperidine.
 - e) (S) 1-Boc-2-(3-Nitro-5-trifluoromethyl-phenoxymethyl)pyrrolidine
 - f) 1-Boc-3-(5-nitro-2-pentafluoroethyl-phenoxymethyl) azetidine.
 - g) N-Boc-[2-(5-nitro-2-pentafluoroethyl-phenoxy)ethyl]amine.
 - h) (R) 3-(2-tert-Butyl-5-nitro-phenoxymethyl)-1-Bocpyrrolidine.
- 25 i) 3-(2-tert-Butvl-5-nitro-phenoxymethyl)-1-Boc-azetidine.
 - j) (S)-1-Boc-[2-(5-nitro-2-tert-butylphenoxymethyl]pyrrolidine
 - k) (S) 3-(2-tert-Butyl-5-nitro-phenoxymethyl)-1-Bocpyrrolidine.
- 30 1) (R)-1-Boc-[2-(5-nitro-2-tert-butylphenoxymethyl]pyrrolidine

Preparation III - 1-Boc-4-(3-amino-5-trifluoromethylphenoxy)-piperidine

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1-Boc-4-(3-nitro-5-trifluoromethyl-phenoxy)-piperidine (470 mg) was dissolved in MeOH (12 ml) and Pd/C (10 mg) was added. After sparging briefly with H₂, the mixture was stirred under H₂ for 6 H. The catalyst was removed by filtration and the MeOH solution was concentrated in vacuo to yield 1-Boc-4-(3-amino-5-trifluoromethyl-phenoxy)-piperidine as an off-white foam.

The following compounds were prepared similarly to the procedure outlined above:

- a) 1-Boc-2-(3-Amino-5-trifluoromethyl-phenoxymethyl)pyrrolidine.
- b) 2-(3-Amino-5-trifluoromethyl-phenoxymethyl)-1-methylpyrrolidine.
 - c) [2-(1-Methylpiperidin-4-yloxy)-pyridin-4-yl]methylamine. ESI (M+H)=222.
 - d) [2-(2-Morpholin-4-vl-ethoxy)-pyridin-4-vl]methylamine.
 - e) [2-(2-Morpholin-4-yl-propoxy)-pyridin-4-yl]methylamine.
 - f) [2-(1-Methyl-pyrrolidin-2-ylmethoxy)-pyridin-4-yl]methylamine. ESI MS: (M+H)=222.
 - g) (4-Aminomethyl-pyridin-2-yl)-(3-morpholin-4-yl-propyl)amine. ESI MS: (M+H)=251.
 - h) 4-tert-Butyl-3-(1-methyl-piperidin-4-ylmethoxy)phenylamine.
 - i) 4-tert-Butyl-3-(2-piperidin-1-yl-ethoxy)-phenylamine.
 - j) 3-(1-Methyl-piperidin-4-ylmethoxy)-4-pentafluoroethylphenylamine.
- k) 3-(1-Isopropyl-piperidin-4-ylmethoxy)-4-pentafluoroethyl-30 phenylamine.
 - 1) (S) 3-Oxiranylmethoxy-4-pentafluoroethyl-phenylamine.
 - m) 3-(2-Pyrrolidin-1-yl-ethoxy)-4-trifluoromethylphenylamine.

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- n) 3-(2-Piperidin-1-yl-ethoxy)-4-trifluoromethylphenylamine.
- o) (S) 3-(1-Boc-pyrrolidin-2-ylmethoxy)-4-pentafluoroethylphenylamine.
- 5 p) (R) 3-(1-Boc-pyrrolidin-2-ylmethoxy)-4-pentafluoroethylphenylamine.
 - q) (R) 3-(1-Methyl-pyrrolidin-2-ylmethoxy)-5trifluoromethyl-phenylamine.
 - r) (S) 3-(1-Methyl-pyrrolidin-2-ylmethoxy)-5trifluoromethyl-phenylamine
 - s) (R) 3-Oxiranylmethoxy-4-pentafluoroethyl-phenylamine.
 - t) (R) 2-(5-Amino-2-pentafluoroethyl-phenoxy)-1-pyrrolidin-1-vl-ethanol.
 - u) 3-(1-Boc-azetidin-3-ylmethoxy)-4-pentafluoroethylphenylamine.
 - v) 3-(2-(Boc-amino)ethoxy)-4-pentafluoroethyl-phenylamine.
 - w) 6-Amino-2,2-dimethyl-4H-benzo[1,4]oxazin-3-one. M+H 193.2. Calc'd 192.1.
 - x) 2,2,4-Trimethyl-3,4-dihydro-2H-benzo[1,4]oxazin-6-ylamine.
 - y) 1-(6-Amino-2,2-dimethyl-2,3-dihydro-benzo[1,4]oxazin-4yl)-ethanone. M+H 221.4. Calc'd 220.3.
 - z) [2-(1-Benzhydryl-azetidin-3-yloxy)-pyridin-4-yl]methylamine.
- 25 aa) [2-(1-Methyl-piperidin-4-ylmethoxy)-pyridin-4-yl]methylamine. M+H 236.3. Calc'd 235.2.
 - ab) 3-(4-Boc-piperazin-1-ylmethyl)-5-trifluoromethylphenylamine. M+H 360.3.
- ac) 2-Boc-4,4-dimethyl-1,2,3,4-tetrahydro-isoquinolin-7-30 vlamine.
- ad) 3-Morpholin-4-vlmethyl-4-pentafluoroethyl-phenylamine.
 - ae) 3-(4-Methyl-piperazin-1-ylmethyl)-4-pentafluoroethylphenylamine. M+H 410.3. Calc'd 409.4.

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- af) 7-Amino-2-(4-methoxy-benzyl)-4,4-dimethyl-3,4-dihydro-2H-isoguinolin-1-one, M+H 311.1.
- ag) 7-Amino-4,4-dimethyl-3,4-dihydro-2H-isoquinolin-1-one.
- ah) (3-Amino-5-trifluoromethyl-phenyl)-(4-Boc-piperazin-1-yl)-methanone. M+H 374.3: Calc'd 373.
- ai) 3-(4-Boc-Piperazin-1-ylmethyl)-5-trifluoromethylphenylamine.
- aj) 1-(7-Amino-4,4-dimethyl-3,4-dihydro-1H-isoquinolin-2-yl)-ethanone. M+H 219.2.
- 10 ak) {2-[2-(1-Methylpiperidin-4-yl)ethoxy]-pyridin-4-yl}methylamine.
 - al) {2-[2-(1-Pyrrolidinyl)ethoxy]-pyridin-4-yl}-methylamine.
 - am) {2-[2-(1-Methylpyrrolin-2-yl)ethoxy]-pyridin-4-yl}methylamine.
- 15 an) (2-Chloro-pyrimidin-4-yl)-methylamine.
 - ao) 3-(1-Boc-azetidin-3-ylmethoxy)-5-trifluoromethylphenylamine.
 - ap) 4-tert-Butyl-3-(1-Boc-pyrrolidin-3-ylmethoxy)phenylamine. M+H 385.
- 20 aq) 4-tert-Butyl-3-(1-Boc-azetidin-3-ylmethoxy)-phenylamine. M+Na 357.
 - ar) (S) 4-tert-Butyl-3-(1-Boc-pyrrolidin-2-ylmethoxy)phenylamine. M+Na 371.
 - as) 3-tert-Butyl-4-(4-Boc-piperazin-1-yl)-phenylamine
- 25 at) 3-(1-Methyl-piperidin-4-yl)-5-trifluoromethylphenylamine.
 - au) 3,3-Dimethyl-2,3-dihydro-benzofuran-6-ylamine.
 - av) 3,9,9-Trimethyl-2,3,4,4a,9,9a-hexahydro-1H-3-aza-fluoren-6-ylamine.
- 30 aw) 4-[1-Methyl-1-(1-methyl-piperidin-4-yl)-ethyl]phenylamine was prepared using EtOH as the solvent.
 - ax) 4-tert-Butyl-3-(4-pyrrolidin-1-yl-but-1-enyl)phenylamine.

- ay) (R) 3-(1-Boc-pyrrolidin-2-ylmethoxy)-5-trifluoromethylphenylamine.
- az) (S) 3-(1-Boc-pyrrolidin-2-vlmethoxy)-5-trifluoromethylphenylamine.

Preparation IV - 1-Boc-4-{3-[(2-fluoro-pyridine-3-carbonyl)aminol-5-trifluoromethyl-phenoxyl-piperidine

1-Boc-4-(3-amino-5-trifluoromethyl-phenoxy)-piperidine (4.37 g) was dissolved in CH₂Cl₂ (100 ml) and NaHCO₁ (2.4 g, Baker)

10 was added. 2-Fluoropyridine-3-carbonyl chloride (2.12 g) was added an the reaction was stirred at RT for 2.5 h. The reaction was filtered and concentrated in vacuo to vield a vellow foam. (30%) EtOAc/Hexane was added and 1-Boc-4-{3-[(2-fluoro-pyridine-3-carbonyl)-amino]-5-trifluoromethyl-

15 phenoxy}-piperidine precipitated as an off white solid.

The following compounds were prepared similarly to the procedure outlined above:

- 20 a) 2-Fluoro-N-[3-(3-piperidin-1-vl-propyl)-5trifluoromethyl-phenyl]-nicotinamide.
 - b) N-[4-tert-Butyl-3-(2-piperidin-1-yl-ethoxy)-phenyl]-2fluoro-nicotinamide.
 - c) N-(3,3-Dimethyl-1-(1-methyl-piperidin-4-ylmethyl)-2,3dihydro-1H-indol-6-yl]-2-fluoro-nicotinamide.
 - d) N-[1-(2-Dimethylamino-acetyl)-3,3-dimethyl-2,3-dihydro-1H-indol-6-yl]-2-fluoro-nicotinamide
 - e) N-[3.3-Dimethyl-1-(2-(Boc-amino)acetyl)-2.3-dihydro-1Hindol-6-yl]-2-fluoro-nicotinamide.
- 30 f) N-(4-Acetyl-2,2-dimethyl-3,4-dihydro-2H-benzo[1,4]oxazin-6-v1)-2-fluoro-nicotinamide, M+H 344.5. Calc'd 343.4.
 - a) 2-Fluoro-N-(2.2.4-trimethyl-3.4-dihydro-2Hbenzo[1,4]oxazin-6-yl)-nicotinamide. M+H 316.2. Calc'd 315.1.

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h) N-(2,2-Dimethyl-3-oxo-3,4-dihydro-2H-benzo[1,4]oxazin-6-yl)-2-fluoro-nicotinamide. M+H 316.1. Calc'd 315.10.

- i) 2-Fluoro-N-[3-(4-methyl-piperazin-1-ylmethyl)-5trifluoromethyl-phenyl]-nicotinamide. M+H 481. Calc'd 480.
- j) 2-Fluoro-N-(2-Boc-4,4-dimethyl-1,2,3,4-tetrahydroisoguinolin-7-yl)-nicotinamide. M+H 400.
- k) 2-Fluoro-N-[3-(4-methyl-piperazin-1-ylmethyl)-4pentafluoroethyl-phenyl]-nicotinamide. M+H 447.0. Calc'd 446.
- 2-Fluoro-N-(3-morpholin-4-ylmethyl-4-pentafluoroethylphenyl)-nicotinamide.
- m) 2-Fluoro-N-[4-iodophenyl]-nicotinamide.
- n) 2-Fluoro-N-(4,4-dimethyl-1-oxo-1,2,3,4-tetrahydroisoquinolin-7-yl)-nicotinamide. M+H 314.0, Calc'd 311.
 - o) 2-Fluoro-N-[3-(4-Boc-piperazine-1-carbonyl)-5trifluoromethyl-phenyl]-nicotinamide. M+H 495.
 - p) 2-Fluoro-N-[3-(4-Boc-piperazin-1-ylmethyl)-5trifluoromethyl-phenyl]-nicotinamide. M+H 483.3; Calc'd 482.
 - q) N-(2-Acetyl-4,4-dimethyl-1,2,3,4-tetrahydro-isoquinolin-7-yl)-2-fluoro-nicotinamide. M+H 430.0.
 - r) N-[3,3-Dimethyl-1-(1-methyl-piperidin-4-yl)-2,3-dihydro-1H-indol-6-yl]-2-fluoro-nicotinamide. M+H 383.2; Calc'd 382.5.
 - s) N-(4-tert-Butylphenyl)-2-fluoronicotinamide.
 - t) N-(4-Trifluoromethylphenyl)-2-fluoronicotinamide.
 - u) 2-Fluoro-N-[3-(1-Boc-azetidin-3-ylmethoxy)-5trifluoromethyl-phenyl]-nicotinamide. M-H 468.2; Calc'd 469.16.
 - v) 2-Fluoro-N-[3-(1-Boc-azetidin-3-ylmethoxy)-4-tert-butyl-phenyl]-nicotinamide.
 - w) (S) N-[4-tert-Butyl-3-(1-Boc-pyrrolidin-2-ylmethoxy)phenyl]-2-fluoro-nicotinamide. M+Na 494.

- x) N-[3-(1-Methyl-piperidin-4-yl)-5-trifluoromethyl-phenyl]-2-fluoro-nicotinamide was prepared with K₂CO₃. instead of NaHCO.
- y) N-(3-Bromo-5-trifluoromethyl-phenyl)-2-fluoronicotinamide.
- z) 2-Fluoro-N-(3,9,9-trimethyl-2,3,4,4a,9,9a-hexahydro-1H-3-aza-fluoren-6-yl)-nicotinamide.
- aa) 2-Fluoro-N-{4-[1-methyl-1-(1-methyl-piperidin-4-yl)ethyl]-phenyl}-nicotinamide
- 10 ab) N-[3,3-Dimethyl-1-(1-Boc-piperidin-4-ylmethyl)-2,3-dihydro-1H-indol-6-yl)-2-fluoro-nicotinamide.

Preparation V - 1-Boc-4-{3-[(2-chloro-pyridine-3-carbonyl)-amino]-5-trifluoromethyl-phenoxy}-piperidine

- 15 1-Boc-4-(3-[(2-chloro-pyridine-3-carbonyl)-amino]-5trifluoromethyl-phenoxy)-piperidine was prepared from 1-Boc4-(3-amino-5-trifluoromethyl-phenoxy)-piperidine and 2chloropyridine-3-carbonyl chloride by a procedure similar to
 that described in the preparation of 1-Boc-4-(3-[(2-fluoro-
- 20 pyridine-3-carbonyl)-amino]-5-trifluoromethyl-phenoxy)piperidine.

The following compounds were prepared similarly to the procedure outlined above:

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- a) N-(4-tert-Butyl-3-nitro-phenyl)-2-chloro-nicotinamide.
- b) 2-Chloro-N-[3-(3-piperidin-1-yl-propyl)-5trifluoromethyl-phenyl]-nicotinamide.
- c) 2-Chloro-N-[3-(3-morpholin-4-yl-propyl)-5-30 trifluoromethyl-phenyl]-nicotinamide.
 - d) 2-Chloro-N-[3-(1-methylpiperidin-4-yl)-5-trifluoromethylphenyl)-nicotinamide.
 - e) 2-Chloro-N-[3-(1-methyl-piperidin-4-ylmethoxy)-4pentafluoroethyl-phenyl]-nicotinamide.

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f) 2-Chloro-N-[3-(1-isopropyl-piperidin-4-ylmethoxy)-4-pentafluoroethyl-phenyll-nicotinamide.

- g) (S) 2-Chloro-N-[4-(oxiranylmethoxy)-3-pentafluoroethylphenyl]-nicotinamide.
- 5 h) 2-Chloro-N-[3-(2-pyrrolidin-1-yl-ethoxy)-4trifluoromethyl-phenyl]-nicotinamide.
 - i) 2-Chloro-N-[3-(2-piperidin-1-yl-ethoxy)-4-pentafluoroethyl-phenyll-nicotinamide.
- j) (R) 2-Chloro-N-[3-(1-Boc-pyrrolidin-2-ylmethoxy)-4-10 pentafluoroethyl-phenyll-nicotinamide.
 - k) (S) 2-Chloro-N-[3-(1-Boc-pyrrolidin-2-ylmethoxy)-4-pentafluoroethyl-phenyl]-nicotinamide.
 - (R) 2-Chloro-N-[3-(1-methyl-pyrrolidin-2-ylmethoxy)-5trifluoromethyl-phenyl]-nicotinamide.
- 15 m) (S) 2-Chloro-N-[3-(1-methyl-pyrrolidin-2-ylmethoxy)-5trifluoromethyl-phenyl]-nicotinamide.
 - n) (R) 2-Chloro-N-[4-(oxiranylmethoxy)-3-pentafluoroethylphenyll-nicotinamide.
- o) (R) Acetic acid 2-(5-[(2-chloro-pyridine-3-carbonyl) amino]-2-pentafluoroethyl-phenoxy)-1-pyrrolidin-1-yl-ethyl ester.
 - p) 2-Chloro-N-[3-(4-methyl-piperazin-1-ylmethyl)-5trifluoromethyl-phenyll-nicotinamide.
 - q) 2-Chloro-N-[2-(4-methoxy-benzyl)-4,4-dimethyl-1-oxo-1,2,3,4-tetrahydro-isoquinolin-7-yl]-nicotinamide. M+H 450.2. Calc'd 449.
 - r) 2-Chloro-N-(4,4-dimethyl-1-oxo-1,2,3,4-tetrahydroisoquinolin-7-yl)-nicotinamide. M+H 330.1, Calc'd 329.
 - s) 2-Chloro-N-[3-(4-Boc-piperazin-1-ylmethyl)-5trifluoromethyl-phenyl]-nicotinamide.
 - t) 2-{3-[(2-Chloro-pyridine-3-carbonyl)-amino]-phenyl}-2methyl-propionic acid methyl ester. M+H 405
 - u) N-{4-tert-Butyl-3-[2-(1-Boc-piperidin-4-yl)-ethyl]phenyl}-2-chloro-nicotinamide. M+Na 524. Calc'd 501.1.

- v) N-[3,3-Dimethyl-1,1-dioxo-2,3-dihydro-1Hbenzo[d]isothiazol-6-vl]-2-chloro-nicotinamide.
- w) N-[1,1,4,4-Tetramethyl-1,2,3,4-tetrahydro-naphth-6-yl]-2chloro-nicotinamide.
- 5 x) 2-Chloro-N-[3,3-dimethyl-2,3-dihydro-benzofuran-6-yl]-2-chloro-nicotinamide.
 - y) 2-Chloro-N-[3-(1-Boc-piperidin-4-yloxy)-5trifluoromethyl-phenyl]-nicotinamide.
 - z) 2-Chloro-N-[3-(1-methyl-piperidin-4-ylmethyl)-5trifluoromethyl-phenyl]-nicotinamide.
 - aa) 2-Chloro-N-[3-(3-piperidin-1-yl-propyl)-5trifluoromethyl-phenyl]-nicotinamide.
 - ab) N-[4-tert-Butyl-3-(4-pyrrolidin-1-yl-but-1-enyl)phenyl]-2-chloro-nicotinamide.
- 15 ac) (R) 2-Chloro-N-[3-(1-Boc-pyrrolidin-2-ylmethoxy)-5-trifluoromethyl-phenyl]-nicotinamide.
 - ad) (S) 2-Chloro-N-[3-(1-Boc-pyrrolidin-2-ylmethoxy)-5trifluoromethyl-phenyl]-nicotinamide.
- 20 Preparation VI 1-Boc-2-(3-[(2-fluoro-pyridine-3-carbonyl)-amino]-5-trifluoromethyl-phenoxymethyl)-pyrrolidine
 - 1-Boc-2-{3-[(2-Fluoro-pyridine-3-carbonyl)-amino]-5trifluoromethyl-phenoxymethyl}-pyrrolidine was prepared from
 1-Boc-2-(3-amino-5-trifluoromethyl-phenoxymethyl)-
- 25 pyrrolidine by a procedure similar to that described in the preparation of 1-Boc-4-(3-[(2-fluoro-pyridine-3-carbonyl)amino]-5-trifluoromethyl-phenoxy)-piperidine.

Preparation VII - 2-(3-nitro-5-trifluoromethyl-

30 phenoxymethyl)-pyrrolidine

1-Boc-2-(3-nitro-5-trifluoromethyl-phenoxymethyl)- pyrrolidine (2.35 g) was dissolved in CH_2Cl_2 (60 ml) and TFA (20 ml) was added. After stirring for 1 h at RT, the mixture was concentrated in vacuo to yield 2-(3-nitro-5-

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trifluoromethyl-phenoxymethyl)-pyrrolidine as an oil that solidified upon standing. The material was used as is without further purification.

- 5 The following compounds were prepared similarly to the procedure outlined above:
 - a) (4-Aminomethyl-pyrimidin-2-yl)-(3-morpholin-4-yl-propyl)amine.
- 10 b) (4-Aminomethyl-pyrimidin-2-yl)-[2-(1-methyl-pyrrolidin-2-yl)-ethyl]-amine.

Preparation VIII - 1-methyl-2-(3-nitro-5-trifluoromethyl-phenoxymethyl)-pyrrolidine

- 2-(3-Nitro-5-trifluoromethyl-phenoxymethyl)-pyrrolidine (6 mmol) was dissolved in CH₃CN (20 ml) and formaldehyde (2.4 ml, 37% aqueous) was added. NaBH₃CN (607 mg) was added, an exotherm was observed. The pH is monitored every 15 min and adjusted to ~7 with AcOH. After 45 min, the mixture was
- 20 concentrated in vacuo and the residue is dissolved in EtOAc, washed with 6N NaOH, 1N NaOH, and 2N HCl (3x). The acid washings were combined, adjusted to -pH 10 with solid Na₂CO₃ and extracted with EtOAc (2x). The EtOAc fractions were combined, dried with Na₂SO₄, and purified with flash
- 25 chromatography (SiO₂, 95:5:0.5 CH₂Cl₂:MeOH:NH₄OH) to afford 1-methyl-2-(3-nitro-5-trifluoromethyl-phenoxymethyl)pyrrolidine.

The following compounds were prepared similarly to the 30 procedure outlined above:

- a) 2-(1-Methylpiperidin-4-yl)-ethanol.
- b) 2-{3-[(2-Fluoro-pyridine-3-carbony1)-amino]-5trifluoromethy1-phenoxymethy1)-1-methylpyrrolidine.

Preparation IX - 4-tert-butyl-3-nitro-phenylamine

A mixture of 1,3-dinitro-4-tert-butylbenzene (10.0~g) in H_2O (56~ml) was heated to reflux. A mixture of Na_2S (21.42~g) and sulfur (2.85~g) in H_2O (34~ml) was added over 1 h via an addition funnel. The reaction maintained at reflux for 1.5 h then cooled to RT and extracted with EtOAc. The organic extracts were combined and washed with H_2O , brine, dried over MgSO₄ and concentrated in vacuo to afford 4-tert-butyl-3-nitro-phenylamine which was used as is without further purification.

Preparation X - N-(3-bromo-5-trifluoromethyl-phenyl)-acetamide

3-Bromo-5-(trifluoromethyl)phenylamine (5 g, Alfa-Aesar) was dissolved in AcOH (140 ml) and Ac₂O (5.9 ml, Aldrich) was added. The reaction was stirred at RT overnight. The mixture was added slowly to H₂O (~700 ml) forming a white precipitate. The solid was isolated by filtration, washed with H₂O and dried under vacuum to yield N-(3-bromo-5-trifluoromethyl-phenyl)-acetamide.

Preparation XI - N-[3-(3-piperidin-1-yl-propyl)-5trifluoromethyl-phenyl]-acetamide

25 Allylpiperidine (1.96 g, Lancaster) was degassed under vacuum, dissolved in 0.5 M 9-BBN in THF (31.2 ml, Aldrich), and heated to reflux for 1 h, then cooled to RT. PD(dppf)Cl₂/CH₂Cl₂ was added to a degassed mixture of N-(3-bromo-5-trifluoromethyl-phenyl)-acetamide, K₂CO₃ (9.8 g) DMF 30 (32.1 ml and H₂O (3 ml). The allyl piperidine solution was added heated to 60°C for 3 h. After cooling to RT and reheating at 60°C for 6 h, the mixture was cooled to RT and poured into H₂O. The mixture was extracted with EtOAc (2x), and the EtOAc portion was washed with 2 N HCl (2x) and

brine. The aqueous phases were combined and the pH was adjusted to ~11 with NaOH (15%) forming a cloudy suspension. The cloudy suspension was extracted with EtOAc (2x) and the EtOAc portion was dried with Na₂SO₄, filtered and concentrated in vacuo. The crude material was purified by flash chromatography (SiO₂, 95:5:0.5 CH₂Cl₂:MeOH:NH₄OH) to afford N-[3-(3-piperidin-1-yl-propyl)-5-trifluoromethyl-phenyl]-acetamide as a brown oil that solidified under vacuum.

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The following compounds were prepared similarly to the procedure outlined above:

- a) N-(3-Morpholin-4-ylpropyl-5-trifluoromethyl-phenyl) acetamide from 4-allyl-morpholine.
- b) N-(3-(1-methylpiperdin-4-ylmethyl-5-trifluoromethyl-phenyl)-acetamide from 1-Methyl-4-methylene-piperidine.

Preparation XII - 3-(3-piperidin-1-yl-propyl)-5-20 trifluoromethyl-phenylamine

N-[3-(3-Piperidin-1-yl-propyl)-5-trifluoromethyl-phenyl]acetamide (1.33 g) was dissolved in EtOH (40 ml) and 12 N
HCl (40 ml) was added. After stirring overnight at 70°C and
RT, the mixture was concentrated in vacuo, affording 3-(3piperidin-1-yl-propyl)-5-trifluoromethyl-phenylamine as a
brown oil.

The following compounds were prepared similarly to the procedure outlined above:

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- a) 3,3-Dimethyl-6-nitro-2,3-dihydro-1H-indole. M+H 193.1; Calc'd 192.2.
- b) 3-(1-Methyl-piperidin-4-ylmethyl)-5-trifluoromethylphenylamine.

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c) 3-Morpholin-4-ylmethyl-5-trifluoromethyl-phenylamine.

Preparation XIII - 3,3-Dimethyl-6-nitro-1-piperidin-4vlmethyl-2,3-dihydro-1H-indole

- 5 3,3-Dimethyl-1-(1-Boc-piperidin-4-ylmethyl)-6-nitro-2,3-dihydro-1H-indole was dissolved in HCl/EtOAc and stirred for 2 h. The mixture was concentrated in vacuo and partitioned between 1,2-dichloroethane and 1N NaOH. The organic layer was removed, washed with brine, dried (Na₂SO₄) and filtered.
 10 The material was used without further purification.
 - Preparation XIV N-[3-(3-morpholin-4-yl-propyl)-5trifluoromethyl-phenyl]-acetamide

N-[3-(3-Morpholin-4-yl-propyl)-5-trifluoromethyl-phenyl]acetamide was prepared from allyl morpholine and N-(3-bromo5-trifluoromethyl-phenyl)-acetamide similar to that
described in the preparation of N-[3-(3-piperidin-1-ylpropyl)-5-trifluoromethyl-phenyl]-acetamide.

- 20 Preparation XV 3-(3-morpholin-4-yl-propyl)-5trifluoromethyl-phenylamine
 - 3-(3-Morpholin-4-yl-propyl)-5-trifluoromethyl-phenylamine was prepared from N-[3-(3-morpholin-4-yl-propyl)-5-trifluoromethyl-phenyl]-acetamide similar to that described in the preparation of 3-(3-piperidin-1-yl-propyl)-5-trifluoromethyl-phenylamine.

Preparation XVI - 1-methyl-4-methylene-piperidine

Ph₃PCH₃I (50 g, Aldrich) was suspended in Et₂O (20 ml) and

30 butyllithium (77.3 ml, 1.6 M in hexanes, Aldrich) was added dropwise. The reaction was stirred for 2 h at RT then 1methylpiperidone (12.3 ml, Aldrich) was added slowly. The mixture was stirred at RT overnight. The solid was removed by filtration, the volume was reduced to -400 ml and

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additional solid was removed by filtration. The Et_2O was washed with H_2O (2x) and 2N HCl (4x). The pH of the acid washings was adjusted to ~11 with 6 N NaOH, then they were extracted with CH_2Cl_2 (4x). The CH_2Cl_2 washings were dried over Na_2SO_4 and concentrated cold in vacuo to provide 1-methyl-4-methylene-piperidine which was used as is.

Preparation XVII - N-[3-(1-methylpiperidin-4-y1)-5-trifluoromethyl-phenyl]-acetamide

N-[3-(1-Methylpiperidin-4-yl)-5-trifluoromethyl-phenyl]acetamide was prepared from 1-methyl-4-methylene-piperidine and N-(3-bromo-5-trifluoromethyl-phenyl)-acetamide similar to that described in the preparation of N-[3-(3-piperidin-1-yl-propyl)-5-trifluoromethyl-phenyl]-acetamide.

Preparation XVIII - 3-(1-methylpiperidin-4-yl)-5trifluoromethyl-phenylamine

3-(1-Methylpiperidin-4-yl)-5-trifluoromethyl-phenylamine was prepared from N-[3-(1-methylpiperidin-4-yl)-5-trifluoromethyl-phenyl)-acetamide similar to the procedure

described in the preparation of 3-(3-piperidin-1-yl-propyl)-5-trifluoromethyl-phenylamine.

Preparation XIX - 2-(1-methylpiperidin-4-yloxy)-4pyridylcarbonitrile

4-Hydroxy-1-methylpiperidine (25.4 g) was dissolved in THF (50 ml) in a 100 mL r.b. flask. NaH/mineral oil mixture (9.58 g) was slowly added to the flask and stirred for 20 min. 2-Chloro-4-cyanopyridine was added to the mixture and stirred at RT until completion. Diluted mixture with EtOAc and added H₂O to quench mixture, then transferred contents to a sep. funnel. The organic phase was collected while the aqueous phase was washed two times with EtOAc. The combined

organics were dried over Na2SO4, filtered, then concentrated

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in vacuo. Then redissolved mixture in CH_2Cl_2 , 10% HCl (300 ml) was added and the mixture was transferred to sep. funnel. The org. was extracted, while EtOAc along with 300 mL 5N NaOH was added to the sep. funnel. The organic phases were collected, dried over Na_2SO_4 , filtered and concentrated in vacuo affording 2-(1-methylpiperidin-4-yloxy)-4-pyridylcarbonitrile as a brown solid. ESI (M+H) = 218.

The following compounds were prepared similarly to the 10 procedure outlined above:

- a) 2-(1-methylpiperidin-4-ylmethoxy)-4-pyridylcarbonitrile. M+H 232.1. Calc'd 231.1.
- b) 2-(1-Benzhydryl-azetidin-3-yloxy)-4-pyridylcarbonitrile.
 15 M+H 342.2. Calc'd 341.2.
 - c) 2-(1-methylpiperidin-4-ylethoxy)-4-pyridylcarbonitrile.
 - d) 2-(1-pyrrolidinylethoxy)-4-pyridylcarbonitrile.
 - e) 2-(1-methylpyrrolin-2-ylethoxy)-4-pyridylcarbonitrile.
 - f) 2-[2-(1-Boc-azetidin-3-yl)-ethoxy]-4-pyridylcarbonitrile.

Preparation XX - [2-(1-methylpiperidin-4-yloxy)-pyridin-4-yl]methylamine bis hydrochloride

[2-(1-Methylpiperidin-4-yloxy)-pyridin-4-yl]methylamine was diluted with Et_2O (50 ml) and 1M $HC1/Et_2O$ (47 ml) was added.

25 The vessel was swirled until precipitate formed.

Preparation XXI - 2-(2-morpholin-4-yl-ethoxy)-4pyridylcarbonitrile

2-(2-Morpholin-4-yl-ethoxy)-4-pyridylcarbonitrile was

30 prepared from 2-chloro-4-cyanopyridine and 2-morpholin-4-ylethanol by a procedure similar to that described in the
preparation of 2-(1-methylpiperidin-4-yloxy)-4pyridylcarbonitrile. The hydrochloride salt was prepared

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similar to that described for [2-(1-methylpiperidin-4yloxy)-pyridin-4-yl]methylamine bis hydrochloride.

Preparation XXII - 2-morpholin-4-yl-propanol

- 5 LAH powder (1.6 g) was added to a flask while under N₂ atmosphere, immediately followed by THF (50 ml). The mixture was chilled to 0°C, methyl 2-morpholin-4-yl-propionate (5 g) was added dropwise to the reaction mixture and stirred at 0°C. After 1 h, the mixture was worked up by adding H₂O (44 mL), 2N NaOH (44 mL), then H₂O (44 mL, 3x). After 30 min of stirring, the mixture was filtered through Celite® and the organic portion was concentrated in vacuo providing 2-morpholin-4-yl-propanol as a colorless oil.
- 15 The following compounds were prepared similarly to the procedure outlined above:
 - a) (1-Methyl-piperidin-4-yl)-methanol. M+H 130.2. Calc'd 129.1.

Preparation XXIII - 2-(2-morpholin-4-yl-propожу)-4pyridylcarbonitrile

2-(2-Morpholin-4-yl-propoxy)-4-pyridylcarbonitrile was prepared from 2-chloro-4-cyanopyridine and 2-morpholin-4-yl-25 propanol by a procedure similar to that described in the preparation of 2-(1-methylpiperidin-4-yloxy)-4pyridylcarbonitrile.

Preparation XXIV - 2-(1-Methyl-pyrrolidin-2-ylmethoxy)-4pyridylcarbonitrile

2-(1-Methyl-pyrrolidin-2-ylmethoxy)-4-pyridylcarbonitrile was prepared from 2-chloro-4-cyanopyridine and 1-methyl-pyrrolidin-2-ylmethanol by a procedure similar to that

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described in the preparation of 2-(1-methylpiperidin-4-vloxy)-4-pyridylcarbonitrile. ESI MS: (M+H)=218.

Preparation XXV - 2-(3-morpholin-4-y1-propylamino)-4pyridylcarbonitrile

To a flask charged with 2-chloro-4-cyanopyridine (2.0 g), was added the aminopropyl morpholine (2.11 ml). The mixture was heated to 79°C for 5 h and stirred. After 5 h the reaction was incomplete. The mixture was then heated at 60°C overnight. The crude compound was purified on silica gel (1-5% MeOH/CH₂Cl₂ gradient). ESI MS: (M+H)=247, (M-H)=245.

Preparation XXVI - 5-Nitro-2-pentafluoroethylphenol

Combined 2-methoxy-4-nitro-1-pentafluoroethylbenzene (9.35 g) and pyridine hydrochloride in a round bottom flask and heated at 210°C for 1 h then cooled to RT. The mixture was diluted with EtOAc and 2N HCl (>500 ml) until all residue dissolved. The organic layer was removed, washed with 2N HCl (2x) and concentrated in vacuo. The residue was dissolved in hexanes and Et₂O, washed with 2N HCl, then brine. Dried organic layer over Na₂SO₄, filtered, concentrated in vacuo and dried under high vacuum to provide 5-nitro-2-pentafluoromethylphenol.

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Preparation XXVII - 2-tert-Butyl-5-nitro-aniline

To $\rm H_2SO_4$ (98%, 389 mL) in a 500 mL 3-neck flask was added 2-tert-butyl aniline (40.6 mL). The reaction was cooled to -10°C and KNO₃ in 3.89 g aliquots was added every 6 min for a total of 10 aliquots. Tried to maintain temperature at -5°C to -10°C. After final addition of KNO₃, stirred the reaction for five min then it was poured onto ice (50 g). The black mix was diluted with $\rm H_2O$ and extracted with EtOAc. The aqueous layer was basified with solid NaOH slowly then

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extracted with EtOAc (2x). The combined organic layers were washed with 6N NaOH and then with a mix of 6N NaOH and brine, dried over Na₂SO₄, filtered and concentrated in vacuo to obtain crude 2-tert-butyl-5-nitro-aniline as a dark red-black oil which solidified when standing at RT. The crude material was triturated with about 130 mL hexanes. After decanting the hexanes, the material was dried to obtain a dark-red black solid.

10 Preparation XXVIII - 2-tert-Butyl-5-nitrophenol

In a 250 ml round bottom flask, 20 mL concentrated $\rm H_2SO4$ was added to 2-tert-butyl-5-nitro-aniline (7.15 g) by adding 5 mL aliquots of acid and sonicating with occasional heating until all of the starting aniline went into solution. $\rm H_2O$

- 15 (84 ml) was added with stirring, then the reaction was cooled to 0°C forming a yellow-orange suspension. A solution of NaNO₂ (2.792 g) in H₂O (11.2 mL) was added dropwise to the suspension and stirred for 5 min. Excess NaNO₂ was neutralized with urea, then the cloudy solution
- 20 was transferred to 500 ml 3-necked round bottom flask then added 17 mL of 1:2 H₂SO₄:H₂O solution, and heated at reflux. Two additional 5 mL aliquots of 1:2 H₂SO₄:H₂O solution, a 7 mL aliquot of 1:2 H₂SO₄:H₂O solution and another 10 mL of 1:2 H₂SO₄: H₂O were added while heating at reflux. The mixture
- 25 was cooled to RT forming a black layer floating on top of the aqueous layer. The black layer was diluted with EtOAc (300 mL) and separated. The organic layer was washed with H₂O then brine, dried over Na₂SO₄ and concentrated in vacuo. Crude oil was purified on silica gel column with 8%
- 30 EtOAc/Hexanes. Upon drying under vacuum, the 2-tert-butyl-5-nitrophenol was isolated as a brown solid.

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Pr paration XXIX - 1-methylpiperidine-4-carboxylic acid thyl ester

Piperidine-4-carboxylic acid ethyl ester (78 g) was dissolved in MeOH (1.2 L) at RT then formaldehyde (37%, 90 ml) and acetic acid (42 ml) were added and stirred for 2 h. The mixture was cooled to 0°C, NaCNBH, (70 g) was added, and the mix was stirred for 20 min at 0°C, then overnight at RT. The mixture was cooled to 0°C then quenched with 6N NaOH. The mixture was concentrated in vacuo to an aqueous layer,

which was extracted with EtOAc (4x), brine-washed, dried over Na₂SO₄, and concentrated in vacuo to provide 1methylpiperidine-4-carboxylic acid ethyl ester.

The following compounds were prepared similarly to the 15 procedure outlined above:

a) (1-Methyl-piperidin-4-yl)-methanol. M+H 130.2. Calc'd 129.1.

20 Preparation XXX - N-[4-tert-Butyl-3-(1-methyl-piperidin-4-ylmethoxy)-phenyl]-2-chloro-nicotinamide

N-[4-tert-Butyl-3-(1-methyl-piperidin-4-ylmethoxy)-phenyl]2-chloro-nicotinamide was prepared from 4-tert-butyl-3-(1methyl-piperidin-4-ylmethoxy)-phenylamine by a procedure

25 similar to that described in the preparation of 1-Boc-4-{3[(2-chloro-pyridine-3-carbonyl)-amino]-5-trifluoromethylphenoxy)-piperidine.

Preparation XXXI - 1-[2-(2-tert-Buty1-5-nitro-phenoxy)-ethyl]-piperidine

To 2-tert-butyl-5-nitrophenol (1.01 g) and K_2CO_3 (1.72 g) was added acetone (35 ml) and H_2O (10.5 mL), then 1-(2-chloroethyl)piperidine HCl (1.909 g) and TBAI (153 mg). The mixture was stirred at reflux overnight. Additional K_2CO_3

(850 mg) and 1-(2-chloroethyl)-piperidine HCl (950 mg) were added and the mixture was heated at reflux for 6 h. The mixture was concentrated in vacuo to an aqueous layer which was acidified with 2N HCl and extracted with EtOAc. The aqueous layer was basified with 6N NaOH and washed with CH_3Cl_2 (3x). The combined organic layers were washed with

CH₂Cl₂ (3x). The combined organic layers were washed with brine/1N NaOH and dried over Na₂SO₄. Washed the EtOAc layer with 2N NaOH/brine and dried over Na₂SO₄. The crude material was purified by silica gel column chromatography

10 with 15% EtoAc/Hexanes to yield 1-[2-(2-tert-butyl-5-nitro-phenoxy)-ethyl]-piperidine as a light tan solid. (M+1)=307.3.

Preparation XXXII - 1-Boc-Piperidine-4-carboxylic acid ethyl

15 ester

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To a stirred solution of piperidine-4-carboxylic acid ethyl ester (23.5 g) in EtOAc (118 ml) at 0°C was added dropwise Boc_2O in EtOAc (60 ml). The reaction was warmed to RT and stirred overnight. Washed reaction with H_2O , 0.1N HCl, H_2O ,

NaHCO₃ and brine. The organic layer was dried over Na₂SO₄, filtered and concentrated in vacuo. The liquid was dried under vacuum to provide 1-Boc-piperidine-4-carboxylic acid ethyl ester.

- 25 The following compounds were prepared similarly to the procedure outlined above:
 - a) N-Boc-(2-chloropyrimidin-4-yl)-methylamine.
 - b) 1-(2-tert-Butyl-4-nitrophenyl)-4-Boc-piperazine.
- 30 c) 1-Boc-azetidine-3-carboxylic acid
 - d) 1-Boc-4-Hydroxymethyl-piperidine using TEA.

Preparation XXXIII - 1-Boc-4-hydroxymethyl-piperidine

1-Boc-4-Hydroxymethyl-piperidine was prepared from 1-Bocpiperidine-4-carboxylic acid ethyl ester by a procedure similar to that described in the preparation of 2-morpholin-

5 4-yl-propanol.

Preparation XXXIV - 1-Boc-4-Methylsulfonyloxymethylpiperidine

Dissolved 1-Boc-4-hydroxymethyl-piperidine in anhydrous

10 CH₂Cl₂ (50 mL) and TEA (4.5 mL) and cooled to 0°C. Mesyl chloride (840 µL) was added and the mixture was stirred for 15 min then at RT for 45 min. The mixture was washed with brine/IN HCl and then brine, dried over Na₂SO₄, concentrated in vacuo and dried under high vacuum to provide 1-Boc-4
15 methylsulfonyloxymethyl-piperidine as a yellow orange thick

5 methylsulfonyloxymethyl-piperidine as a yellow orange thic} oil.

The following compounds were prepared similarly to the procedure outlined above:

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a) 1-Boc-3-methylsulfonvloxymethyl-azetidine.

Preparation XXXV - 1-Boc-4-(3-nitro-6-pentafluoroethylphenoxymethyl)-piperidine

25 To a slurry of 60% NaH suspension in DMF (30 mL) at RT added a solution of 5-nitro-2-pentafluoroethyl-phenol (3.6 g) in 5 mL DMF. The dark red mixture was stirred at RT for 10 min then added a solution of 1-Boc-4-methylsulfonyloxymethyl-piperidine (3.1 g) in 5 mL DMF. The reaction was stirred at 30 60°C and 95°C. After 1h, added 2.94 g K₂CO₃ and stirred overnight at 105°C. After cooling to RT, the reaction was diluted with hexanes and 1N NaOH. Separated layers, and washed organic layer with 1N NaOH and with brine, dried over

Na2SO4, filtered and concentrated in vacuo. Purification

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with silica gel column chromatography with 8% EtOAc/Hexanes yielded 1-Boc-4-(3-nitro-6-pentafluoroethyl-phenoxymethyl)piperidine as a light yellow thick oil.

5 Preparation XXXVI - 4-(3-nitro-6-pentafluoroethylphenoxymethyl)-piperidine

4-(3-Nitro-6-pentafluoroethyl-phenoxymethyl)-piperidine was prepared from 1-Boc-4-(3-nitro-6-pentafluoroethylphenoxymethyl)-piperidine by a procedure similar to that

10 described in the preparation of 2-(3-nitro-5trifluoromethyl-phenoxymethyl)-pyrrolidine.

Preparation XXXVII - 1-methyl-4-(3-nitro-6-pentafluoroethyl-phenoxymethyl)-piperidine

15 4-(3-Nitro-6-pentafluoroethyl-phenoxymethyl)-piperidine (316.5 mg) was dissolved in 2.7 mL acetonitrile, then added 37% formaldehyde/H₂O (360 ul) and then NaBH₂CN (90 mg). Upon addition of NaCNBH3 the reaction exothermed slightly. The reaction was stirred at RT and pH was maintained at ~7 2.0 by addition of drops of glacial acetic acid. After about 1 h, the mixture was concentrated in vacuo, treated with 8 mL 2N KOH and extracted two times with 10 mL Et20. The organic layers were washed with 0.5N KOH and then the combined organic layers were extracted two times with 1N HCl. The 25 aqueous layer was basified with solid KOH and extracted two times with Et20. This organic layer was then washed with brine/1N NaOH, dried over Na2SO4, filtered, concentrated in

30 Preparation XXXVIII - 1-Isopropyl-4-(5-nitro-2pentafluoroethyl-phenoxymethyl)-piperidine

Dissolved 4-(5-nitro-2-pentafluoroethyl-phenoxymethyl)piperidine (646 mg) in 1,2-dichloroethane (6.4 ml), then added acetone (136 ul), NaBH(OAc)₃ (541 mg) and finally

vacuo and dried under high vacuum to give pure compound.

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acetic acid (105 ul). Stirred the cloudy yellow solution under N_2 at RT overnight. Added another 130 uL acetone and stirred at RT over weekend. Quenched the reaction with 30 mL N NaOH/H₂O and stirred 10 min. Extracted with Et₂O and the organic layer was brine-washed, dried over Na₂SO₄, filtered and concentrated in vacuo. Dried under high vacuum for several h to obtain 1-isopropyl-4-(5-nitro-2-pentafluoroethyl-phenoxymethyl)-piperidine as a yellow orange solid.

10 The following compounds were prepared similarly to the

procedure outlined above:

- a) 3,3-Dimethyl-1-(1-methyl-piperidin-4-yl)-6-nitro-2,3dihydro-1H-indole was prepared using 1-methyl-piperidin-4-one. M+H 290: Calc'd 289.4.
- b) 3,3-Dimethyl-1-(1-Boc-piperidin-4-ylmethyl)-6-nitro-2,3dihydro-1H-indole using 1-Boc-4-formyl-piperidine.

Preparation XXXIX - 3,3-Dimethyl-1-(1-methyl-piperidin-4-20 ylmethyl)-6-nitro-2,3-dihydro-1H-indole

3,3-Dimethyl-1-piperidin-4-ylmethyl-6-nitro-2,3-dihydro-1H-indole was treated with an excess of formaldehyde and NaBH(OAc)₃ and stirred overnight at RT. The reaction was quenched with MeOH and concentrated in vacuo. The residue was partitioned between EtOAc and 1N NaOH. The organic layer was removed, washed with brine, dried (Na₂SO₄), filtered and

Preparation XL - (S) 2-(5-Nitro-2-pentafluoroethyl-

concentrated to provide the compound.

30 phenoxymethyl)-oxirane

Combined 5-nitro-2-pentafluoromethylphenol (2.69 g), DMF (25 ml) K_2CO_3 (3.03 g) and (S) toluene-4-sulfonic acid oxiranylmethyl ester (2.27 g) and stirred the mixture at 90°C. After about 4 hours, the mix was cooled, diluted with EtOAc,

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washed with H_2O , 1N NaOH (2x), 1N HCl and then with brine. Dried over Na_2SO_4 , filtered and concentrated in vacuo. Purified the crude on silica gel column with 5% EtOAc/hexane and drying under high vacuum provided the (S)-2-(5-nitro-2-pentafluoroethyl-phenoxymethyl)-oxirane.

The following compounds were prepared similarly to the procedure outlined above:

a) (R)-2-(5-Nitro-2-pentafluoroethyl-phenoxymethyl)-oxirane.

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Preparation XLI - (S) 2-Chloro-N-[3-(2-hydroxy-3-pyrrolidin-1-y1-propoxy)-4-pentafluoroethy1-pheny1]-nicotinamide

(S) 2-Chloro-N-[4-(2-oxiranylmethoxy-)-3-pentafluoroethyl-phenyl]-nicotinamide (1.11 g) in a sealed tube and added pyrrolidine (285 μl). Stirred after sealing tube at 60°C. After 12 h, the mix was concentrated in vacuo and purified on a silica gel column (5:95:0.5 MeOH:CH₂Cl₂:NH₄OH - 8:92:1, MeOH:CH₂Cl₂:NH₄OH). Concentrated in vacuo and dried under high vacuum to obtain pure compound.

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The following compounds were prepared similarly to the procedure outlined above:

a) (R) 1-(5-Nitro-2-pentafluoroethyl-phenoxy)-3-pyrrolidin-1-vl-propan-2-ol.

Preparation XLII - 5-nitro-2-trifluoromethylanisole

Cooled 140 mL pyridine in a large sealable vessel to -40° C. Bubbled in trifluoromethyl iodide from a gas cylinder which had been kept in freezer overnight. After adding ICF3 for 20 min, added 2-iodo-5-nitroanisole (24.63 g) and copper powder (67.25 g). Sealed vessel and stirred vigorously for 22 h at 140°C After cooling to -50°C, carefully unsealed reaction vessel and poured onto ice and Et20. Repeatedly

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washed with Et₂O and H_2O . Allowed the ice - Et₂O mixture to warm to RT. Separated layers, washed organic layer with 1N HCl (3x), then brine, dried over Na₂SO₄, filtered and concentrated in vacuo. Eluted material through silica gel plug (4.5:1 Hex:CH₂Cl₂) to provide 5-nitro-2-trifluoromethylanisole.

Preparation XLIII - 1-[2-(5-nitro-2-trifluoromethylphenoxy)ethyl]pyrrolidine

10 1-[2-(5-Nitro-2-trifluoromethylphenoxy)ethyl]-pyrrolidine was prepared from 5-nitro-2-trifluoromethyl-phenol and 1-(2chloroethyl)pyrrolidine by a procedure similar to that described for 1-[2-(2-tert-butyl-5-nitro-phenoxy)-ethyl]piperidine.

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Preparation XLIV - 1-[2-(5-Nitro-2-pentafluoroethyl-phenoxy)-ethyl]-piperidine

1-[2-(5-Nitro-2-pentafluoroethyl-phenoxy)-ethyl]-piperidine
was prepared from 5-nitro-2-pentafluoroethylphenol and 1-(220 chloroethyl)piperidine by a procedure similar to that
described in the preparation of 1-[2-(2-tert-butyl-5-nitrophenoxy)-ethyl]-piperidine.

Preparation XLV - 3-(1-Boc-pyrrolidin-2-ylmethoxy)-4-

25 pentafluoroethyl-phenylamine

3-(2-Pyrrolidin-1-yl-methoxy)-4-trifluoromethyl-phenylamine was prepared from 1-[2-(5-nitro-2-trifluoromethylphenoxy)methyl]-pyrrolidine by a procedure similar to that described in the preparation of 1-Boc-4-(3-

30 amino-5-trifluoromethyl-phenoxy)-piperidine.

Preparation XLVI - 2-Chloro-N-[3-(2-pyrrolidin-1-yl- thoxy)-4-trifluorom thyl-phenyl]-nicotinamide

2-Chloro-N-[3-(2-pyrrolidin-1-yl-ethoxy)-4-trifluoromethyl-phenyl]-nicotinamide was prepared from 3-(2-pyrrolidin-1-yl-ethoxy)-4-trifluoromethyl-phenylamine and 2-chloropyridine-3-carbonyl chloride by a procedure similar to that described in the preparation of 1-Boc-4-(3-[(2-chloro-pyridine-3-carbonyl)-amino]-5-trifluoromethyl-phenoxyl-piperidine.

10 Preparation XLVII - (R) Acetic acid 2-(5-nitro-2pentafluoroethyl-phenoxy)-1-pyrrolidin-1-ylmethyl-ethyl ester

Dissolved 1-(5-nitro-2-pentafluoroethyl-phenoxy)-3-pyrrolidin-1-yl-propan-2-ol (3.5 g) in CH₂Cl₂ (15 ml), added TEA (2.55 ml) and cooled to 0°C. Acetyl chloride (781.3 µl) was added dropwise, forming a suspension. The mixture was warmed to RT and stirred for 1.5 h. Additional acetyl chloride (200 µl) was added and the mix was stirred for another h. The mixture was diluted with CH₂Cl₂ and washed with sat. NaHCO₃. The organic layer was removed, washed with brine and back extracted with CH₂Cl₂. Dried the combined organic layers over Na₂SO₄, filtered and concentrated in vacuo. The residue was purified over silica gel column (5:94.5:0.5 MeOH: CH₂Cl₂:NH₄OH) to provide acetic acid 2-(5-nitro-2-pentafluoroethyl-phenoxy)-1-pyrrolidin-1-

The following compounds were prepared similarly to the procedure outlined above:

ylmethyl-ethyl ester as a yellow brown oil.

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- a) (R) Acetic acid 2-(5-amino-2-pentafluoroethyl-phenoxy)-1pyrrolidin-1-yl-methyl-ethyl ester.
- b) 1-(2,2-Dimethyl-6-nitro-2,3-dihydro-benzo[1,4]oxazin-4-yl)-ethanone. M-NO₂ 206.4; Calc'd 250.1.

Preparation XLVIII - (R) 2-Chloro-N-[3-(2-hydroxy-2pyrrolidin-1-yl-propoxy)-4-pentafluoroethyl-phenyl]nicotinamide

5 (R) Acetic acid 2-(5-[(2-chloro-pyridine-3-carbonyl)-amino]2-pentafluoroethyl-phenoxy}-1-pyrrolidin-1-yl-ethyl ester
(408 mg) was dissolved in MeOH (15 ml) and NH4OH (6 ml) was
added and the mixture was stirred at RT for 6 h. The
reaction was concentrated in vacuo and dried under high
10 vacuum. The residue was purified over silica gel column
(8:92:0.6 MeOH: CH2Cl2:NH4OH). The purified fractions were
concentrated in vacuo and dried again to provide (R)-2chloro-N-[3-(2-hydroxy-2-pyrrolidin-1-yl-ethoxy)-4pentafluoroethyl-phenyl|-nicotinamide as a white foam.

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Preparation XLIX - 2-Dimethylamino-1-(3,3-dimethyl-6-nitro-2,3-dihydro-indol-1-yl)-ethanone

3,3-Dimethyl-6-nitro-2,3-dihydro-1H-indole (5 g) was dissolved in DMF (100 ml) and HOAt (3.89 g) dimethylamino20 acetic acid (5.83 g) and EDC (3.89 g) were added. The reaction was stirred overnight. The mixture was diluted with CH₂Cl₂ (1L) and washed with sat'd NaHCO₃ (3x200 ml). The organic layer was washed with brine, dried over Na₂SO₄, filtered and concentrated in vacuo. The residue was purified by flash chromatography (SiO₂, EtOAc to 5%MeOH/EtOAc) to afford the title compound.

The following compounds were prepared similarly to the procedure outlined above:

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a) 1-(3,3-Dimethyl-6-nitro-2,3-dihydro-indol-1-yl)-2-(N-Boc-amino)-ethanone.

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Preparation L - 1-(6-Amino-3,3-dimethyl-2,3-dihydro-indol-1-yl)-2-(N-Boc-amino)-ethanone

- 1-(3,3-Dimethyl-6-nitro-2,3-dihydro-indol-1-yl)-2-(N-Bocamino)-ethanone (3.9 g) was dissolved in EtOH (30 ml) and Fe powder (3.1 g) NH₄Cl (299 mg) and H₂O (5 ml) were added. The reaction was stirred at 80°C overnight. The reaction was filtered through Celite® and evaporated off the MeOH. The residue was partitioned between CH₂Cl₂ and sat'd NaHCO₃. The organic layer was removed, washed with brine, dried over
- 10 Na_2SO_4 , filtered and concentrated in vacuo. The residue was purified by flash chromatography (SiO_2 , 25% EtOAc/hexane). The purified fractions were concentrated in vacuo to afford the compound as a white powder.
- 15 The following compounds were prepared similarly to the procedure outlined above:
 - a) 1-(6-Amino-3,3-dimethyl-2,3-dihydro-indol-1-yl)-2dimethylamino-ethanone.
- 20 b) 3,3-Dimethyl-1-(1-methyl-piperidin-4-ylmethyl)-2,3-dihydro-1H-indol-6-ylamine.
 - c) 3-(4-Methyl-piperazin-1-ylmethyl)-4-pentafluoroethylphenylamine. M+H 324.2. Calc'd 323.
 - d) 3,3-Dimethyl-1-(1-methyl-piperidin-4-yl)-2,3-dihydro-1Hindol-6-vlamine, M+H 259.6: Calc'd 259.3.
 - e) 3,3-Dimethyl-1,1-dioxo-2,3-dihydro-1H-116benzo[d]isothiazol-6-ylamine
 - f) 1,1,4,4-Tetramethyl-1,2,3,4-tetrahydro-naphth-6-ylamine.
- g) 3,3-Dimethyl-1-(1-Boc-piperidin-4-ylmethyl)-2,3-dihydro-30 1H-indol-6-vlamine.

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Preparation LI - 2-Boc-4,4-dimethyl-7-nitro-1,2,3,4-tetrahydro-isomuinoline

4,4-Dimethyl-7-nitro-1,2,3,4-tetrahydro-isoquinoline (150 mg) was dissolved with CH₂Cl₂ (3 ml) DIEA (100 ul) DMAP (208 mg and Boc₂O (204 mg) and the mixture was stirred for 6 h at RT. The reaction was diluted with CH₂Cl₂, washed with sat'd NaHCO₃ and dried over MgSO₄, filtered and concentrated to provide the compound which was used without further purification.

The following compounds were prepared similarly to the procedure outlined above substituting Ac₂O:

15 a) 1-(4,4-Dimethyl-7-nitro-3,4-dihydro-1H-isoquinolin-2-yl)ethanone. M+H 249.3.

Preparation LII - 2-Bromo-N-(4-methoxy-benzyl)-5-nitrobenzamide

20 PMB-amine (5.35 ml) in CH₂Cl₂ (130 ml) was slowly added to 2-bromo-5-nitro-benzoyl chloride (10.55 g) and NaHCO₃ (9.6 g) and the mixture was stirred at RT for 1 h. The mixture was diluted with CH₂Cl₂ (1 L), filtered, washed with dilute HCl, dried, filtered again, concentrated and dried under 25 vacuum to provide the compound as a white solid. M+H 367.

Calc'd 366.

Preparation LIII - 2-Bromo-N-(4-methoxy-benzyl)-N-(2-methyl-allyl)-5-nitro-benzamide

30 To a suspension of NaH (1.22 g) in DMF (130 ml) was added 2-bromo-N-(4-methoxy-benzyl)-5-nitro-benzamide (6.2 g) in DMF (60 ml) at -78C. The mixture was warmed to 0°C, 3-bromo-2-methyl-propene (4.57 g) was added and the mixture was stirred for 2 h at 0°C. The reaction was poured into ice

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water, extracted with EtOAc (2x400 ml), dried over MgSO₄, filtered and concentrated to a DMF solution which was used without further purification.

5 Preparation LIV - of 2-(4-Methoxy-benzyl)-4,4-dimethyl-7nitro-3,4-dihydro-2H-isoguinolin-1-one

2-Bromo-N-(4-methoxy-benzyl)-N-(2-methyl-allyl)-5-nitrobenzamide (23.4 mmol) was dissolved in DMF (150 ml) and $\rm Et_4NC1$ (4.25 g), $\rm HCO_2Na$ (1.75 g) and $\rm NaOAc$ (4.99 g) were

10 added. N_2 was bubbled through the solution for 10 min, then $Pd(OAc)_2$ (490 mg) was added and the mixture was stirred overnight at $70^{\circ}C$. The mixture was extracted with EtOAc, washed with sat'd NH_4C1 , dried over $MgSO_4$, filtered and concentrated until the compound precipitated as a white 15 solid.

The following compounds were prepared similarly to the procedure outlined above:

- 20 a) 3,3-Dimethyl-6-nitro-2,3-dihydro-benzofuran was prepared from 1-bromo-2-(2-methyl-allyloxy)-4-nitro-benzene.
 - b) 3,9,9-Trimethyl-6-nitro-4,9-dihydro-3H-3-aza-fluorene was prepared from 4-[1-(2-bromo-4-nitro-phenyl)-1-methylethyl]-1-methyl-1,2,3,6-tetrahydro-pyridine.

Preparation LV - 4,4-Dimethyl-7-nitro-3,4-dihydro-2Hisoquinolin-1-one

2-(4-Methoxy-benzyl)-4,4-dimethyl-7-nitro-3,4-dihydro-2H-isoquinolin-1-one (2.0 g) was dissolved in CH_3CN (100 ml) and H_2O (50 ml) and cooled to $0^{\circ}C$. CAN (9.64 g) was added and the reaction was stirred at $0^{\circ}C$ for 30 min, then warmed to RT and stirred for 6 h. The mixture was extracted with CH_2Cl_2 (2x300 ml) washed with sat'd NH_4Cl , dried over $MgSO_4$, filtered and concentrated. The crude material was

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recrystallized in CH₂Cl₂/EtOAc (1:1) to give 4,4-dimethyl-7-nitro-3,4-dihydro-2H-isoquinolin-1-one as a white solid.

Preparation LVI - 4,4-Dimethyl-7-nitro-1,2,3,4-tetrahydroisoguinoline

4,4-Dimethyl-7-nitro-3,4-dihydro-2H-isoquinolin-1-one (230 mg) was dissolved in THF (10 ml) and BH_3Me_2S (400 ul) was added and the reaction was stirred overnight at RT. The reaction was quenched with MeOH (10 ml) and NaOH (200 mg)

- and heating at reflux for 20 min. The mixture was extracted with EtOAc, washed with sat'd NH₄Cl, extracted with 10% HCl (20 ml). The acidic solution was treated with 5N NaOH (15 ml), extracted with EtOAc (30 ml) dried, filtered and evaporated to give the compound as a yellow solid. M+H
- 15 207.2, Calc'd 206.

The following compounds were prepared similarly to the procedure outlined above:

20 a) 4-Boc-2,2-dimethyl-6-nitro-3,4-dihydro-2Hbenzo[1,4]oxazine.

Preparation LVII - 2-Bromomethyl-4-nitro-1-pentafluoroethylbenzene

- 25 2-Methyl-4-nitro-1-pentafluoroethyl-benzene (2.55 g) was dissolved in CCl₄ (30 ml) and AIBN (164 mg) and NBS (1.96 g) were added. The reaction was heated to reflux and stirred for 24 h. The mix was diluted with CH₂Cl₂, washed with sat'd NaHCO₃, dried over MgSO₄ and concentrated to give the
- 30 compound as an oil which was used without further purification.

Preparation LVIII - 1-Methyl-4-(5-nitro-2-pentafluoro thylbenzyl)-piperazine

2-Bromomethyl-4-nitro-1-pentafluoroethyl-benzene (2.6 g) was added to N-methylpiperazine (5 ml) and stirred at RT for 3 h. The mixture was filtered and the filtrate was treated with 1-chlorobutane, extracted with 5N NaOH (6 ml). The acidic solution was treated with 5N NaOH (6 ml) then extracted with EtOAc. The organic layer was removed, dried over MgSO₄ and concentrated to give the compound as an oil.

The following compounds were prepared similarly to the procedure outlined above:

15 a) 4-(5-Nitro-2-pentafluoroethyl-benzyl)-morpholine.

Preparation LIX - 1-Boc-4-(5-nitro-2-pentafluoroethyl-benzyl)-piperazine.

20 2-Bromomethyl-4-nitro-1-pentafluoroethyl-benzene (2.5 g) was dissolved in CH₂Cl₂ and added to N-Boc-piperazine (2.5 g) and NaHCO₃ (1 g) and stirred at RT overnight. The mixture was diluted with CH₂Cl₂ (100 ml), washed with sat'd NH₄Cl, dried over MgSO₄, filtered and concentrated. The residue was purified by silica gel chromatography (hexane, CH₂Cl₂:hexane 2:8) to give the compound as an yellow solid.

Preparation LX - (4-Boc-piperazin-1-yl)-(3-nitro-5trifluoromethyl-phenyl)-methanone

30 A mixture of 3-nitro-5-trifluoromethyl-benzoic acid (4.13 g), 4-Boc-piperazine (2.97 g), EDC (3.88 g), HOBt (2.74 g), DIEA (3.33 ml) in CH₂Cl₂ (120 ml) was stirred at RT for 3 h. The mixture was diluted with CH₂Cl₂ (100 ml), washed with sat'd NH₄Cl₁, dried over MgSO₄, filtered and concentrated.

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The residue was purified by silica gel chromatography (hexane, CH_2Cl_2 :hexane 1:2) to give the compound as a white solid.

5 Preparation LXI - 1-Boc-4-(3-nitro-5-trifluoromethylbenzyl)-piperazine

(4-Boc-piperazin-1-y1)-(3-nitro-5-trifluoromethy1-pheny1)methanone (403 mg) was dissolved in THF (6 ml) and BH₃Me₂S
(300 µ1) was added and the reaction was stirred for 3 h at
60°C and 2 h at RT. The reaction was quenched with MeOH (5
ml) and NaOH (100 mg) and stirred at RT for 1 h. The
mixture was concentrated and dissolved in CH₂Cl₂, washed
with sat'd NH₄Cl/NaHCO₁, dried (MgSO₄), filtered and

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Preparation LXII - 2-Ethyl-4-aminomethyl pyridine

evaporated to give the compound as an oil. M+H 390.3.

To a solution of 2-ethyl-4-thiopyridylamide (10 g) in MeOH (250 ml) was added Raney 2800 Nickel (5 g, Aldrich) in one portion. The mixture was stirred at RT for 2 days then at 60°C for 16 h. The mixture was filtered, concentrated to provide the desired compound.

Preparation LXIII - N-Boc-[2-(4-morpholin-4-yl-butyl)pyrimidin-4-ylmethyl]-amine

- N-Boc-(2-chloropyrimidine)-methylamine (663 mg) and 4-(aminopropyl)morpholine (786 mg) were dissolved in MeOH and concentrated in vacuo. The residue was heated at 100°C for 15 min, forming a solid which was dissolved in CH₂Cl₂/MeOH then concentrated again and heated 15 min more.
- 30 Concentrated in vacuo and dried under high vacuum. Triturated with a small amount of IpOH and allowed to settle over a weekend. Filtered, rinsing with a small amount of IpOH to provide the compound as a white solid.

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The following compounds were prepared similarly to the procedure outlined above:

a) (4-Bocaminomethyl-pyrimidin-2-yl)-[2-(1-methylbyrrolidin-2-yl)-ethyl]-amine. M+H 336.5; Calc'd 335.45.

Preparation LXIV - 2-fluoronicotinic acid

In a flame dried 3-necked round bottom flask equipped with a dropping funnel and thermometer, under N2, THF (250 ml) was added via cannula. LDA (2M in cyclohexane, 54 ml) was added via cannula as the flask was cooled to -78°C. At -78°C, 2fluoropyridine (8.87 ml) was added dropwise over 10 min. The reaction was stirred for 3 h. Condensation was blown off (with N2) a few cubes of solid CO2 and they were added to the mixture. The mixture was warmed to RT once the solution turned yellow, and it was stirred overnight. The reaction was cooled to 0°C and the pH was adjusted to ~2.5 with 5N HCl. The mixture was concentrated in vacuo and extracted with EtOAc. The EtOAc layer was washed with brine, dried over MgSO4, filtered and concentrated to dryness. The resulting solid was slurried in EtOAc (100 ml), filtered. washed with cold EtOAc and dried at 50°C for 1 h to afford 2-fluoronictinic acid. M+H 142.1; Calc'd 141.0.

25 Preparation LXV - 4-cyano-2-methoxypyridine

added to MeOH (36 ml) with a considerable exotherm. After the Na is dissolved, a solution of 2-chloro-4-cyanopyridine (15 g) in dioxane:MeOH (1:1, 110 ml) was added via dropping 30 funnel over a 10 min period. The reaction was heated to reflux for 3.5 h then cooled at ~10°C overnight. Solid was filtered off and the solid was washed with MeOH. The filtrate was concentrated to ~60 ml and H₂O (60 ml) was added to redissolve a precipitate. Upon further

Under a stream of N2 and with cooling, Na metal (2.7 g) was

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concentration, a precipitate formed which was washed with H₂O. Further concentration produced additional solids. The solids were combined and dried in vacuo overnight at 35°C to provide 4-cvano-2-methoxypyridine which was used as is.

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Preparation LXVI - (2-methoxypyridin-4-y1)methylamine 4-Cyano-2-methoxypyridine (1.7 g) was dissolved in MeOH (50 ml) and conc. HCl (4.96 ml) was added. Pd/C (10%) was added and H $_2$ was added and let stand overnight. The solids were filtered through Celite and the cake was washed with MeOH (~250 ml). Concentration in vacuo produced an oil which was dissolved in MeOH (~20 ml). Et $_2$ O (200 ml) was added and

stirred for 1 h. The resulting precipitate was filtered and

15 yl)methylamine (hydrochloride salt) as an off-white solid.

washed with Et₂O to afford (2-methoxypyridin-4-

Preparation LXVII - 2-(4-Amino-pheny1)-2-methyl-propionic acid methyl ester

2-Methyl-2-(4-nitro-phenyl)-propionic acid methyl ester (2.1 g) was dissolved in THF (70 ml) and acetic acid (5 ml) and Zn (10 g) were added. The mixture was stirred for 1 h and filtered through Celite*. The filtrate was rinsed with EtOAc and the organics were evaporated to a residue which was purified on silica gel chromatography (40%EtOAc/hexanes) to provide the desired compound as a yellow oil. M+H 194.

Preparation LXVIII - 1-(2-tert-Butyl-phenyl)-4-methyl-piperazine

2-tert-Butyl-phenylamine and bis-(2-chloro-ethyl)30 methylamine were mixed together with K2CO3 (25 g), NaI (10
g) and diglyme (250 mL) and heated at 170°C for 8 h. Cooled
and filtered solid and evaporated solvent. Diluted with
EtOAc, washed with NaHCO3 solution, extracted twice more

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with EtOAc, washed with brine, dried over Na₂SO₄ and evaporated to give the compound as a dark solid.

The following compounds were prepared similarly to the procedure outlined above:

a) 1-Bromo-2-(2-methyl-allyloxy)-4-nitro-benzene was prepared from methallyl bromide.

Preparation LXIX 3-(1-Methyl-1,2,3,6-tetrahydro-pyridin-4-yl)-5-trifluoromethyl-phenylamine

3-(5,5-Dimethyl-[1,3,2]dioxaborinan-2-yl)-5-trifluoromethylphenylamine (8.8g, 0.032mol)was added to trifluoromethanesulfonic acid 1-methyl-1,2,3,6-tetrahydro-pyridin-4-

15 yl ester (7.91g, 0.032mol)and 2N Na₂CO₃ aqueous solution (25mL) was bubbled through N₂ for 5 min. Pd(PPh₃)₄ (3.7g, 3.2mmol) was added and the reaction was heated to 80°C for 16 h. The reaction was cooled to RT and diluted with Et₂O (100 mL). The mixture was filtered through Celite® and the

20 filtrate was washed with NaHCO₃ aqueous solution (25 ml) followed by brine (25 mL). The organic phase was dried over Na₂SO₄ and concentrated in vacuo. The desired product was isolated by passing through silica gel column chromatography (EtOAc, then (2M NH₃) in MeOH/EtOAc) to provide a yellow

25 oil.

Preparation LXX - 3,3-Dimethyl-6-nitro-2,3-dihydrobenzo[d]isothiazole 1,1-dioxide

3,3-Dimethyl-2,3-dihydro-benzo[d]isothiazole 1,1-dioxide was added to KNO3 in H₂SO₄ cooled to 0°C and stirred for 15 min. The reaction was warmed to RT and stirred overnight. The mix was poured into ice and extracted with EtOAc (3x), washed with H₂O and brine, dried and evaporated to give the product which was used without further purification.

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The following compounds were prepared similarly to the procedure outlined above:

5 a) 1,1,4,4-Tetramethyl-6-nitro-1,2,3,4-tetrahydronaphthalene

Preparation LXXI - 3-(1-Methyl-1,2,3,4-tetrahydro-pyridin-4-vl)-5-trifluoromethyl-phenylamine

- 10 3-(5,5-Dimethyl-[1,3,2]dioxaborinan-2-yl)-5-trifluoromethylphenylamine (1.2 g) was added to trifluoro-methanesulfonic acid 1-methyl-1,2,3,6-tetrahydro-pyridin-4-yl ester (1.0 g), LiCl (500 mg, Aldrich), PPh₃ (300 mg, Aldrich) and 2M Na_2CO_3 aqueous solution (6 ml) and was bubbled with N_2 for 5 min.
- Pd(PPH₃)₄ (300 mg, Aldrich) was added and the reaction was heated to 80°C for 16 h. The reaction was cooled to RT and diluted with Et₂O (100 mL). The mixture was filtered through Celite[®] and the filtrate was washed with NaHCO₃ aqueous solution (25 mL) followed by brine (25 mL). The organic phase was dried over Na₂SO₄ and concentrated in
 - vacuo. The desired compound was isolated by silica gel column chromatography (EtOAc 10% (2M NH₃) in MeOH/EtOAc) to provide yellow oil. M+H 257.2; Calc'd 256.1.

25 Preparation LXXII - Trifluoromethylsulfonic acid 1-methyl-1,2,3,6-tetrahydro-pyridin-4-yl ester

In a three-necked round bottom flask equipped with a thermometer and an additional funnel was placed anhydrous THF (200 mL) and 2M LDA (82.8 mL). The solution was cooled to-78°C and a solution of 1-methyl-piperidin-4-one (20 mL) in anhydrous THF (70 mL) was added drop-wise. The reaction was warmed to -10°C over 30 min and cooled down again to -78°C. Tf₂NPh (54.32 g) in 200 mL of anhydrous THF was added through the additional funnel over 30 min and anhydrous THF

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v1)-5-trifluoromethv1-phenvlamine

(30 mL) was added to rinse the funnel. The reaction was warmed to RT and the reaction solution was concentrated in vacuo. The residue was dissolved in Et_2O purified on neutral Al_2O_3 column chromatography (Et_2O as elutant). The product was obtained as orange oil. (20 g)

Preparation LXXIII - 3-(5,5-Dimethyl-[1,3,2]dioxaborinan-2-

N₂ was bubbled through a solution of 3-bromo-5-

10 trifluoromethyl-phenylamine (2.38 g), 5,5,5',5'-tetramethyl[2,2']bi[[1,3,2]dioxaborinanyl] (2.24 g, Frontier
Scientific) and KOAc (2.92 g), dppf (165 mg, Aldrich) in
anhydrous dioxane (50 ml) for 2 min. PdCl₂ (dppf) (243 mg,
Aldrich) was added and the reaction was heated to 80°C for 4
15 h. After cooling to RT. the mix was diluted with 50 mL of

Et₂O, filtered through Celite*, and the filtrate was concentrated *in vacuo*. The residue was dissolved in Et₂O (100 mL), washed with sat. NaHCO₃ aqueous solution (50 mL) followed by brine (50 mL). The organic phase was dried over Na₂SO₄ and concentrated *in vacuo*. The residue was

dissolved in 3:2 Et₂O/Hex (100 mL), filtered through Celite[®] and the filtrate was concentrated *in vacuo* to afford a dark brown semi-solid.

25 Preparation LXXIV - 1-Boc-3-Hydroxymethyl-azetidine

A solution of 1-Boc-azetidine-3-carboxylic acid (1.6 g) and Et₃N (2 ml) in anhydrous THF (60 ml) was cooled to 0°C. Isopropyl chloroformate (1.3 g) was added via a syringe slowly; forming a white precipitate almost immediately. The reaction was stirred for 1 h at 0°C and the precipitate was filtered out. The filtrate was cooled to 0°C again and aqueous NaBH₄ solution (900 mg, 5 ml) was added via pipette and stirred for 1 h. The reaction was quenched with NaHCO₃ solution (50 mL) and the product was extracted with EtOAc

3.0

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(200 mL). The organic phase was washed with brine (50 mL), dried over Na₂SO₄ and concentrated *in vacuo*. The residue was dissolved in EtOAc and passed through a short silica gel pad. Concentrating the filtrate *in vacuo* provided the compound as a light yellow oil.

Preparation LXXV - 1-Boc-3-(3-nitro-5-trifluoromethyl-phenoxymethyl)-azetidine

A mixture of 1-Boc-3-methylsulfonyloxymethyl-azetidine (1.47 g), 3-nitro-5-trifluoromethyl-phenol (1.15 g) and K₂CO₃ (1.15 g) in DMF(20 ml) at 80°C was stirred overnight. The reaction was cooled to RT and diluted with 25 mL of sat. NaHCO₃ and 50 mL of EtOAc. The organic phase was separated and washed with brine (25 mL), dried over Na₂SO₄ and 15 concentrated *in vacuo*. The crude compound was purified by column chromatography (50% EtOAc/hex).

Preparation LXXVI - 2,2-Dimethyl-6-nitro-3,4-dihydro-2H-benzo[1.4]oxazine

20 2,2-Dimethyl-6-nitro-4H-benzo[1,4]oxazin-3-one was added to BH₃-THF complex (Aldrich) in THF with ice cooling. The mixture was heated to reflux for 2 h then carefully diluted with 12 mL of MeOH and heated to reflux for an additional 1 h. Concentrated HCl (12 mL) was added and heated to reflux for 1 h. The mixture was concentrated and the resulting

solid was suspended in a dilute aqueous solution of NaOH (1 M) and extracted with EtOAc (100 mL x 4). The organic layers were washed with H₂O and dried over MgSO₄. Evaporation of solvent gave a yellow solid.

Preparation LXXVII - 2,2,4-Trimethyl-6-nitro-4H-benzo[1,4]oxazin-3-one

2,2-Dimethyl-6-nitro-4H-benzo[1,4]oxazin-3-one (1.1 g) was mixed with MeI (850 mg, Aldrich), K_2CO_3 (1.38 g, Aldrich)

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and DMF (30 ml, Aldrich) at $40^{\circ}\mathrm{C}$ for 48 h. The DMF was removed in vacuo and the residue was diluted with EtOAc (80 ml). The organic phase was washed with H₂O (50 ml), aqueous Na₂SO₃ (50 ml) and brine (50 ml). The resulting solution was dried (MgSO₄) and concentrated to provide the compound which was used as is

Preparation LXXVIII - 2-Bromo-N-(2-hydroxy-5-nitro-phenyl)-2-methyl-propionamide

- 10 2-Amino-4-nitro-phenol (3.08 g, Aldrich) was stirred with THF (30 ml, Aldrich) in an ice bath. 2-Bromo-2-methyl-propionyl bromide (2.47 ml, Aldrich) and Et₃N (2.0 g, Aldrich) was slowly added via syringe. The mixture was stirred for 45 min then poured into ice. The aqueous phase 15 was extracted by EtOAc (50 mL x 4). The organic layer was dried and concentrated. The desired product was
 - dried and concentrated. The desired product was crystallized from EtOAc. (Chem. Pharm. Bull 1996, 44(1) 103-114).

20 Preparation LXXIX - 2,2-Dimethy1-6-nitro-4H-

benzo[1,4]oxazin-3-one

- 2-Bromo-N-(2-hydroxy-5-nitro-phenyl)-2-methyl-propionamide was mixed with K_2CO_3 in 20 mL of DMF and stirred overnight at $50\,^{\circ}C$. The reaction mixture was poured into ice water.
- 25 The precipitate was collected by filtration and washed with H₂O. The crude compound was recrystallized from EtOH.

Preparation LXXX -4-[1-(2-Bromo-4-nitro-phenyl)-1-methylethyl]-1-methyl-pyridinium iodide

30 l-Methyl-4-[l-methyl-1-(4-nitro-phenyl)-ethyl]-pyridinium (8 g) was dissolved in glacial HOAc (10 ml) then diluted with H₂SO₄ (50 ml), then NBS (3.8 g) was added. After 1 h, additional NBS (1.2 g) was added, 30 min later another 0.5 g of NBS, then 15 min later 200 mg more NBS. After 1 h, the

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mixture was neutralized with NH_4OH (conc.) with ice bath cooling. The neutralized mixture was then concentrated and used as is

- Preparation LXXXI 4-[1-(2-Bromo-4-nitro-phenyl)-1-methylethyl]-1-methyl-1,2,3,6-tetrahydro-pyridine
 - $4-[1-(2-Bromo-4-nitro-phenyl)-1-methyl-ethyl]-1-methyl-pyridiniumiodide was mixed with MeOH (400 ml) and <math>CH_2Cl_2$ (200 ml), then treated with NaBH₄ (2.5 g) in portions.
- After stirring at RT for 2 h, the mixture was extracted with CH₂Cl₂ (300 mL x 3). The CH₂Cl₂ layer was washed with brine, dried over Na₂SO₄ and concentrated in vacuo, to provide the desired product.
- 15 Preparation LXXXII 1-Methyl-4-[1-methyl-1-(4-nitrophenyl)-ethyl1-pyridinium iodide

4-(4-Nitro-benzyl)-pyridine (4.3 g) was mixed with MeI (4 ml, 9.12 g)/NaOH (5N, 30 ml), Bu_4NI (150 mg) and CH_2Cl_2 (50 ml) and stirred at RT overnight. Additional MeI (2 mL) was added along with 50 mL of NaOH (5N). 6 h later, more MeI (2 mL) was added. The mixture was stirred at RT over the weekend. The mixture was cooled on ice bath and the base was neutralized by conc. HCl (aq) addition dropwise to pH 7. The compound was used as is.

Preparation LXXXIII - 1-Methyl-4-(4-nitro-benzyl)-1,2,3,6-tetrahydro-pyridine

4-(4-Nitrobenzyl)pyridine (64 g) and TBAI (6 g) were dissolved in CH₂Cl₂ (500 mL) and the solution was suspended with NaOH (aq. 5N, 450 mL) in a 3L 3-necked round bottom flask. With vigorous stirring, iodomethane (213 g) was added and stirred vigorously at RT for 60 h (or until blue color disappears). The reaction was quenched with dimethylamine (100 mL) and MeOH (300 mL) and stirred for 2 h. NaBH4 (19

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g) was added to the mixture in small portions. The reaction mixture was stirred for 30 min at RT, then partitioned between CH_2Cl_2/H_2O (500 mL/500 mL). The organic layer was collected and the aqueous layer was washed with CH_2Cl_2 (300 mL x 3). The combined organic layers was washed with brine then concentrated in vacuo. The residue was purified on a silica wash-column (7% TEA in EtOAc). The desired fractions were combined and concentrated under vacuum to give the desired compound as a dark gray solid. (MS: M+1=261).

Preparation LXXXIV - 1-Boc-4-formylpiperidine

4A Molecular sieves were heated to 100°C and a vacuum was applied. They were cooled to RT and purged with N2. CH_2Cl_2 (420 ml) and CH_3CN (40 ml), NMO (40 g) and 1-Boc-4-hydroxymethylpiperidine (50 g) were added and the mix was stirred for 5 min then cooled to 15°C . TPAP (4.1 g) is added and an exotherm was observed. The reaction was maintained at RT with external cooling. The reaction was stirred at RT for 3 h, filtered, concentrated, diluted with $50^{\circ}\text{EtOAc/hexanes}$ and purified on a silica gel plug ($50^{\circ}\text{EtOAc/hexanes}$). The eluant fractions were concentrated to afford a yellow oil.

25 Preparation LXXXV 2-Chloro-4-cyanopyridine

2-Chloro-4-cyanopyridine was prepared similar to the method described by Daves et al., J. Het. Chem., 1, 130-32 (1964).

Preparation LXXXVI 4-(2-tert-Butyl-5-nitro-phenyl)-but-3-en1-ol

A mix of 1-(tert-buty1)-2-bromo-4-nitrobenzene (3.652 g), TEA (5.92 ml), 3-buten-1-ol (5.48 ml), $Pd(OAc)_2$ (32 mg), $Pd(PPh_3)_4$ (327 mg) and toluene (40 ml) was degassed with nitrogen and heated in a sealed vessel for 16 h at 120°C.

The next day, the reaction mixture was cooled to RT, filtered, and concentrated in vacuo. The crude was eluted on a silica gel column with 15% to 22% EtOAc/hexanes gradient system to yield a yellow-brown oil.

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Preparation LXXXVII 4-(2-tert-Buty1-5-nitro-pheny1)-but-3-ena1

4-(2-tert-Buty1-5-nitro-pheny1)-but-3-en-1-ol (1.024 g) was dissolved in 10 ml of CH₂Cl₂ and added dropwise over 5 min to a -78C mix of oxalyl chloride (0.645 ml), DMSO (0.583 ml), and 10 ml CH₂Cl₂. The reaction was stirred at -78C for 1 h, then treated with a solution of TEA (1.52 ml) in 7 ml CH₂Cl₂ and stirred at -78C for an additional 25 min, then warmed to -30°C for 35 min. The reaction was treated with 15 50 ml of saturated aqueous NH₄Cl, diluted with H₂O and extracted with EtOAc. The organic layer was brine-washed, dried over Na₂SO₄, filtered, and concentrated in vacuo to yield a yellow oil which was used as is in Preparation

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LXXXVTII.

Preparation LXXXVIII 1-[4-(2-tert-Butyl-5-nitro-phenyl)-but3-enyl]-pyrrolidine

4-(2-tert-Buty1-5-nitro-pheny1)-but-3-enal (895 mg) was dissolved in 40 ml THF, and to the solution was added

- pyrrolidine (0.317 ml). To the deep orange solution was added NaBH(OAc) $_3$ (1.151 g) and glacial AcOH (0.207 ml). The reaction was stirred at RT overnight, then treated with saturated aqueous NaHCO $_3$ and diluted with Et $_2$ O and some 1N NaOH. The layers were separated, and the organic layer was
- 30 extracted with aqueous 2N HCl. The acidic aqueous layer was basified to pH>12 with 6 N NaOH, extracted with Et₂O, brinewashed, dried over Na₂SO₄, filtered, and concentrated in vacuo to provide 1-[4-(2-tert-butyl-5-nitro-phenyl)-but-3enyl]-pyrrolidine as a orange-brown oil.

HOGENOU DIEFOR

Preparation LXXXVIV N-Boc-(2-chloropyrimidin-4-yl)-methylamine

To 2-chloropyrimidine-4-carbonitrile [2.5 g, prepared by the procedure of Daves et. al. [J. Het. Chem. 1964, 1, 130-132)] in EtOH (250 ml) under N₂ was added Boc₂O (7.3 g). After the mixture was briefly placed under high vacuum and flushed with N₂, 10% Pd/C (219 mg) was added. H₂ was bubbled though the mixture (using balloon pressure with a needle outlet) as it stirred 4.2 h at RT. After filtration through Celite⁸, addition of 1.0 g additional Boc₂O, and concentration, the residue was purified by silica gel chromatography (5:1 \rightarrow 4:1 hexanes/EtOAc) to obtain N-Boc-(2-chloropyrimidin-4-yl)-methylamine.

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Example 1

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N-(4-Chloropheny1)(3-[(4-pyridylmethyl)amino](2-thienyl))carboxamide

Step A - Preparation of 3-[(tert-

25 <u>butoxy)carbonylamino]thiophene-2-carboxylic acid</u>
To a mixture of methyl 3-amino-2-thiophenecarboxylate (8 g, 51 mmol) and BOC₂O (11 g, 50 mmol) in CH₂Cl₂ (400 ml) was added 4-(dimethylamino)pyridine (1 g, 8.1 mmol).

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The reaction was stirred at RT overnight and washed with 1N HCl (100 ml), followed by water and brine. The organic layer was dried over Na_2SO_4 and evaporated under reduced pressure and used for the next step without further purification. To the residue (2 g, ~7 mmol) in EtOH (50 ml) was added 1N NaOH (25 ml), the reaction was stirred at RT for 1 h and the solvent was evaporated under reduced pressure. Water (5 ml) was added and the solution was acidified with HOAc. The precipitate was filtered and used in the next step without further purification. MS (ES-): 242 (M-H) $^-$.

Step B - Preparation of {3-[(tert-butoxy)carbonylamino](2-thienyl)}-N-(4-chlorophenyl)carboxamide

To a mixture of the thienyl carboxylic acid from Step A (300 mg, 1.23 mmol) and 4-chloroaniline (160 mg, 1.25 mmol) and DIEA (300 μ l, 1.6 mmol) was added EDC (300 mg, 1.6 mmol) and HOBt (170 mg, 1.25 mmol) in CH₂Cl₂, the reaction was stirred at RT overnight. The solution was washed with 1N HCl and saturated NaHCO₃, followed by H₂O and brine. The organic layer was dried over Na₂SO₄ and evaporated under reduced pressure and purified with preparative TLC to give the amide. MS (ES+): 353 (M+H)*.

25 Step C - Preparation of N-(4-chlorophenyl)(3-[(4-pyridylmethyl)amino](2-thienyl))carboxamide

The amide from Step B was mixed with 25% TFA/CH $_2$ Cl $_2$ and stirred at RT for 1 h (monitored by HPLC). The solvent was evaporated under reduced pressure and the residue was mixed with 4-pyridine carboxaldehyde (260 mg, 2.5 mmol) and NaCNBH $_3$ (160 mg, 2.5 mmol) in MeOH (40 ml). The reaction was stirred at RT overnight and evaporated under reduced pressure. The final product was purified by prep-HPLC as TFA

salt. MS (ES+): 344 (M+H)*; (ES-): 342 (M-H)⁻. Calc'd. for $C_{12}H_{14}C1N_3OS$ - 343.84.

Example 2

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N-Pheny1{3-[(4-pyridylmethy1)amino](2thieny1)}carboxamide

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The title compound was analogously synthesized by method described in Example 1. The final product was purified by preparative HPLC as TFA salt. MS (ES+): 310 (M+H) $^+$; (ES-): 308 (M-H) $^-$. Calc'd. for $C_{17}H_{15}N_5OS$ - 309.4.

Example 3

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N-(4-Chlorophenyl) {2-[(4-pyridylmethyl)amino](3pyridyl)}carboxamide

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Step A - Preparation of (2-amino(3-pyridyl))-N-(4chlorophenyl) carboxamide

To a mixture of 2-aminonicotinic acid (5.3 g, 38 mmol) and 4-chloroaniline (4.9 g, 38 mmol) and DIEA (9 ml, 48 5 mmol) at 0°C in CH2Cl2 was added EDC (9.5 g, 48 mmol) and HOBt (5.1 g, 38 mmol), the reaction was warmed to RT and stirred overnight. The solvent was evaporated under reduced pressure and quenched with 2N NaOH solution (60 ml) and stirred for 20 min. The precipitate was filtered to give the titled compound. MS (ES+): 248 (M+H)+: (ES-): 246 (M-H).

Step B - Preparation of N-(4-chlorophenyl) {2-[(4pyridylmethyl)aminol(3-pyridyl)}carboxamide

To a mixture of the pyridyl carboxamide (400 mg, 1.6 mmol) from Step A and 4-pyridinecarboxaldehyde (200 μ l, 2 mmol) and HOAc (200 µl) in CH2Cl2 was added NaBH(OAc)3 (600 mg, 2.8 mmol), the reaction was stirred at RT overnight. The reaction mixture was washed with H2O and brine and dried over Na2SO4. The solution was evaporated and purified by prep-TLC to give the title compound. MS (ES+): 339 (M+H)+; 20 (ES-): 337 (M-H) -. Calc'd for C18H15ClN4O - 338.796.

Example 4

N-(3,4-Dichlorophenyl) (2-[(4-pyridylmethyl)amino] (3-pyridyl)}-carboxamide

The title compound was analogously synthesized by the method described in Example 3. MS (ES+): 373 (M+H) $^+$; (ES-): 370.9 (M-H) $^-$. Calc'd for $C_{18}H_{14}Cl_2N_4O$ - 373.24.

Example 5

N-(3-Chlorophenyl){2-[(4-pyridylmethyl)amino](3-pyridyl)}carboxamide

The title compound was analogously synthesized by the method described in Example 3. MS (ES+): 339 (M+H) $^+$; (ES-): 337 (M-H) $^-$. Calc'd. for $C_{18}H_{15}ClN_4O$ -338.1.

Example 6

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N-(4-Chlorophenyl) {3-[(4-pyridylmethyl)amino](2-pyridyl)}carboxamid

The title compound was analogously synthesized by 5 method described in Example 3. MS (ES+): 339 (M+H) $^+$; (ES-): 337 (M-H) $^-$. Calc'd. for $C_{18}H_{13}ClN_4O$ - 338.8

Example 7

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HOOFECC DILLOGS

N-(4-Chlorophenyl) {3-[(6-quinolylmethyl)amino](2pyridyl)}carboxamide

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The title compound was analogously synthesized by the method described in Example 3. MS (ES+): 389 (M+H)*; (ES-): 387 (M-H)⁻. Calc'd, for C₂-H₁₇ClN₄O - 388.86.

Example 8

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N-(3,4-Dichlorophenyl) {2-[(6-quinolylmethyl)amino](3pyridyl)}-carboxamide

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The title compound was analogously synthesized by the method described in Example 3. MS (ES+): 423 (M+H)⁺; (ES-): 421 (M-H)⁻. Calc'd. for C₂₂H₁₆Cl₂N₆O - 423.30.

Example 9

N-(4-Chlorophenyl) {6-methyl-2-[(4-pyridylmethyl)amino](3pyridyl)}carboxamide

Step A - Preparation of 6-methyl-2-[(4pyridylmethyl)amino]pyridine-3-carboxylic acid

The mixture of 2-chloro-6-methyl-nicotinic acid (1.0 eq.) and 4-aminomethyl-pyridine (2.0 eq.) was stirred in a sealed tube at 130 °C overnight. The resulted mixture was cooled to RT, diluted with $\rm CH_2Cl_2$, filtered to collected the brown solid. The brown solid was recrystallized in ethanol to give the substituted amine as light brown solid. MS (ES+): 244 (M+H) $^{\circ}$.

Step B - Preparation of N-(4-chlorophenyl) (6-methyl-2-[(4-pyridylmethyl)amino](3-pyridyl))-carboxamide

To the mixture of the substituted amine from Step A (1.0 eq.) and 4-chloroaniline (2.0 eq) in CH_2Cl_2 was added bis(2-oxo-3-oxazolidinyl)phosphinic chloride (1.1 eq.) and TEA (1.1 eq.). The mixture was stirred overnight, diluted with CH_2Cl_2 , washed with saturated NH_4Cl solution, dried over Na_2SO_4 , filtered and concentrated, purified by flash

30 chromatography (4% MeOH/CH₂Cl₂) to give the title compound

as an white solid. MS (ES+): 353 (M+H); (ES-): 351 (M-H). Calc'd. for $C_{19}H_{17}C1N_4O$ - 352.82.

Example 10

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N-(3,4-Dichloropheny1){6-methy1-2-[(4-pyridylmethy1)amino](3-pyridyl)}carboxamide

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TOTAGOL CLASSO

The title compound was analogously synthesized by the method described in Example 9. MS (ES+): 387 (M+H); (ES-): 385 (M-H). Calc'd. for $C_{19}H_{16}Cl_2N_4O$ - 387.27.

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Example 11

N-(3-Fluoro-4-methylphenyl) (6-methyl-2-[(4-pyridylmethyl)amino](3-pyridyl))carboxamide

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The title compound was analogously synthesized by the method described in Example 9. MS (ES+): 351 (M+H); (ES-): 349 (M-H). Calc'd. for C₂₀H₁₉FN₄O - 350.39.

5 Example 12

HICHESEL CHACCE

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(6-Chloro-2-[(4-pyridylmethy1)amino](3-pyridyl)}-N-(4pyridylmethy1)carboxamide

The title compound was analogously synthesized by the method described in Example 9. MS (ES+): 354 (M+H); (ES-): 352 (M-H). Calc'd. for $C_{18}H_{16}C1N_5O$ - 353.81.

Example 13

N-(3,4-Dichloropheny1) {6-chloro-2-[(4-pyridylmethy1)amino](3-pyridyl)}carboxamide

The title compound was analogously synthesized by the method described in Example 9. MS (ES+): 409 (M+H). Calc'd. for C18H18ClNN0 - 407.7.

Example 14

N-(4-Chlorophenyl)(6-chloro-2-[(4-pyridylmethyl)amino](3pyridyl))carboxamide

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HOOFWAY LOTHOUR

The title compound was analogously synthesized by method described in Example 9. MS (ES+): 374 (M+H); (ES-): 372 (M-H). Calc'd. for $C_{18}H_{16}Cl_{2}N_{4}O$ - 373.24.

Example 15

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{6-Chloro-2-[(4-pyridylm thyl)amino](3-pyridyl)}-N-(3fluorophenyl)carboxamide

The title compound was analogously synthesized by the method described in Example 9. MS (ES+): 357 (M+H); (ES-): 355 (M-H). Calc'd. for $C_{18}H_{19}FN_4OC1$ - 356.5.

Example 16

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HOUTERON DATEDON

N-(3-Chlorophenyl) {6-chloro-2-[(4-pyridylmethyl)amino](3pyridyl))carboxamide

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The title compound was analogously synthesized by the method described in Example 9. MS (ES+): 374 (M+H); (ES-): 372 (M-H). Calc'd. for $C_{18}H_{14}Cl_{2}N_{4}O$ - 373.24.

Example 17

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N-(4-Chlorophenyl) [3-[(4-pyridylmethyl ne)amino] (4-pyridinecarboxamide

A mixture of (2-amino(4-pyridyl))-N-(4
5 chlorophenyl)carboxamide (350 mg, 1.4 mmol) (similar procedure to Example 3, Step A) and 4-pyridine carboxaldehyde (200 µl, 2 mmol) and 4-toluenesulfonic acid monohydrate (50 mg) in EtoH (50 ml) was heated to reflux overnight. The solvent was evaporated and the residue was
10 purified by prep-TLC. MS (ES+): 337 (M+H)*; (ES-): 335 (M-H)*. Calc'd. for ChaHnClNO - 336.8.

Example 18

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N-(4-Chlorophenyl) {3-[(4-pyridylmethyl)amino](4-pyridyl)}carboxamide

The compound from Example 17 was mixed with NaBH4 (100 mg) in EtOH (20 ml) and heated to reflux for 5 min. The solvent was evaporated under reduced pressure and the residue was purified by prep-TLC to give the titled compound. MS (ES+): 339 (M+H)*; (ES-): 337 (M-H)*. Calc'd.

25 for CueHisClN40 - 338.8.

Exampl 19

N-(3-Fluoro-4-methylphenyl){2-[(4-pyridylmethyl)amino](3pyridyl)}carboxamide

The title compound was analogously synthesized by the 10 method in Examples 17-18. MS (ES-): 337 (M-H). Calc'd. for $C_{19}H_{17}FN_4O\ -\ 336.37.$

Example 20

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N-(4-Chloropheny1) {2-[(4-quinolylmethy1) amino] (3pyridy1)}carboxamide

ACCHEGA SALSE

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The title compound was analogously synthesized by the method described in Examples 17-18. MS (ES+): 389 (M+H)⁺; (ES-): 387 (M-H)⁻. Calc'd. for C₂₂H₁₂ClN₄O - 388.86.

5 Example 21

N-(4-Chlorophenyl) {2-[(6-quinolylmethyl)amino](3pyridyl)}carboxamide

The title compound was analogously synthesized by the method described in Examples 17-18. MS (ES+): 389 (M+H)*; (ES-): 387 (M-H)⁻. Calc'd. for C₂₂H₁₇ClN₄O - 388.86.

Example 22

20 N-(4-Chlorophenyl)(2-[(4-pyridylethyl)amino]-5-(3-thienyl)-(3-pyridyl))carboxamid

Step A - Preparation of 5-bromo-2-hydroxynicotinic acid

A solution of sodium hypobromide was made by adding Br_2 (1.01 ml, 39.5 mmol, 1.1 eq) slowly over a period of 5 min to NaOH (5N, 40 ml) that was previously cooled to 0°C in an ice bath. The solution was stirred for 10 min before adding 2-hydroxynicotinic acid (5.0 g, 35.9 mmol) and placed in a 50°C oil bath and stirred. Concurrently, a second pot of sodium hypobromide solution was made by slowly adding Br_2 (1.01 ml, 39.5 mmol, 1.1 eq) to a NaOH solution (5N, 40 ml) in an ice bath. The second pot of sodium hypobromide was added to the solution of 2-hydroxynicotinic acid after 24 h of heating then was stirred for an additional 24 h. The solution was cooled to RT, placed in an ice bath and acidified with concentrated HCl while stirring. The precipitate which formed was filtered, washed and dried to afford the desired compound as an off-white solid.

Step B - Preparation of 5-bromo-2-chloronicotinic acid

A solution of 5-bromo-2-hydroxynicotinic acid, from

Step A (8.3 g, 38.1 mmol) and SOCl₂ (40 ml) in a 150 ml
round bottom flask was placed in an 80°C oil bath and
stirred while adding 10 ml of DMF. The solution was heated
at reflux for 4 h at 80°C before cooling to RT. Excess
SOCl₂ was stripped off under reduced pressure forming a

25 yellow-brown residue. The yellow-brown residue was placed
in an ice bath and cooled to 0°C. Residual SOCl₂ was
neutralized and the chloro compound was precipitated by the
dropwise addition of water. Precipitate was filtered,
washed and dried to afford the desired chloro compound as a

light yellow solid.

Step C - Preparation of 5-bromo-2-chloro-N-(4chlorophenyl)nicotinamide

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To a mixture of 4-chloroanaline (594 mg, 4.7 mmol, 1. eq.), EDC (1.62 g, 8.5 mmol, 2 eq.), HOBT (572 mg, 4.2 mmol, 1 eq.), and DIEA (1.1 ml, 6.3 mmol, 1.5 eq.) in CH_2Cl_2 (50 ml) was added 5-bromo-2-chloronicotinic acid from Step B (1.0 g, 4.2 mmol). The reaction was stirred at RT overnight. The solution was quenched with water and the organic layer was purified by chromatography (50% EtOAc in hexane) to afford a light-yellow compound. MS (ES+): 347.0, 349.0 (M+H)*; (ES-): 345.0, 347.0 (M-H) $^{-}$.

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Step D - Preparation of 5-(3-thiophene)-2-chloro-N-(4-chlorophenyl)nicotinamide

3-Thiophene boronic acid (204 mg, 1.6 mmol, 1.1 eq), $Pd(OAC)_2$ (33 mg, 0.2 mmol, 0.2 eq.), and K_2CO_3 (505 mg, 4.3 mmol, 3 eq.) were added to a solution of 5-bromo-2-chloro-N-(4-chlorophenyl)nicotinamide from Step C (500 mg, 1.4 mmol) in DMF (20 ml). The reaction was placed in a 50°C oil bath and stirred overnight. The reaction was filtered and purified by medium pressure chromatography (30% EtOAc in hexane) to afford the desired thienyl compound as an off white solid.

Step E - Preparation of N-(4-chlorophenyl)(2-[(4pyridylethyl)amino]-5-(3-thienyl)-(3-pyridyl))carboxamide.

25 4-(Aminoethyl)pyridine (10 ml) was added to a 25 ml round-bottom flask containing 5-(3-thiophene)-2-chloro-N-(4-chlorophenyl)nicotinamide from Step D (200 mg, 0.6 mmol). The solution was placed in an 80°C oil bath and stirred overnight. The reaction was cooled to RT, and after an 30 aqueous work-up, was purified by medium-pressure chromatography (80% EtOAc in hexane) to afford the title compound as a light yellow solid. MS: (ES+) 435.1 (M+H); (ES-) 432.8 (M-H). Calc'd. for ColHyclNyCS - 434.95.

Example 23

5 N-(4-Chlorophenyl){ 5-(4-methoxyphenyl)-2-[(4-pyridylmethyl)amino]-(3-pyridyl)}carboxamide

The title compound was prepared analogously to Example 22. MS: (ES+) 445.1 (M+H). Calc'd. for $C_{25}H_{21}ClN_4O$ - 444.92.

Example 24

N-(4-Chlorophenyl) (5-bromo-2-[(4-pyridylmethyl)amino]-(3-pyridyl)) carboxamide

The title compound was prepared analogously to Example 22, Steps A, B, C and E. MS: (ES+) 419 (M+H) (ES-) 417 (M-20 H). Calc'd. for $C_{18}H_{14}BrClN_4O$ - 417.69.

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Example 25

N-(4-Isopropylphenyl) {2-[(4-pyridylmethyl)amino](3pyridyl)}carboxamide

Step A: Preparation of (2-chloro-3-pyridyl)-N-(4-isopropylphenyl)carboxamide

To a mixture of 2-chloronicotinic acid (6.3 g) and 4-isopropylaniline (5.26 ml) and DIEA (10 ml) in CH_2Cl_2 (200 ml) was added EDC (10 g) and HOBt (5.4 g). The reaction was stirred at RT overnight and washed with 2 N NaOH (100 ml), H_2O (250 ml) and brine (100 ml). The organic layer was dried over Na_2SO_4 and evaporated to give (2-chloro-3-pyridyl)-N-(4-isopropylphenyl)-carboxamide.

Step B: Preparation of N-[4-(isopropyl)phenyl][2-[(4-pyridylmethyl)amino][3-pyridyl)]carboxamide hydrochloride

A mixture of (2-chloro(3-pyridyl))-N-(4-isopropylphenyl)carboxamide (1.5 g, from Step A) and 4-aminomethylpyridine (0.71 ml) was heated at 130°C neat for 3 h. The reaction was cooled and diluted with $\rm CH_2Cl_2$ and washed with $\rm H_2O$ twice followed by brine. The organic layer was dried with $\rm Na_2SO_4$ and evaporated under reduced pressure. The residue was purified by column chromatography with EtOAc and further mixed with MeOH and 1 N HCl/Et₂O (2 ml). The solution was evaporated to furnish the titled compound. MS

(ES+): 347 (M+H)*; (ES-): 345 (M-H). Calc'd. for $C_{21}H_{22}N_4O$ = 346.18.

The following compounds (Examples 26-81) were synthesized by the method described in Example 25 unless specifically described. Detailed intermediate preparations are included.

Table 1.

	#	Y	R ¹	R ²	M+H	calc'd
	26	-NH-CH ₂ -		Н	347	346.2
15	27	NH-CH ₂ -	P F OH	Н	356	355.1
	28	NH-CH ₂ -	F	н	471	470.1
	29	NH-CH ₂ -		Н	352	351.4
	30	NH-CH ₂ -	N.N.	н	365	364.2

Table 1. cont.

	5	#	Y	R*	R*	М+Н са	lc'd
				N			
j. O		31	NH-CH ₂ -	ji _s »	Н	368	367.5
inguacoi		32	NH-CH ₂ -	N-N-N-N-N-N-N-N-N-N-N-N-N-N-N-N-N-N-N-	н	369	368.5
		33	NH-CH ₂ -	0	н	377	376.2
Company of the Property of the		34	NH-CH ₂ -		Н	366.8	366.8
C N	10	35	NH-CH ₂ -		5-Br	447.0	445.7
		36	NH-CH ₂ -		Н	425.0	424.5
		37	NH-CH ₂ -		Н	363.2	362.4
		38	NH-CH ₂ -		Н	393.2	392.4
		39	NH-CH ₂ -	OH	Н	393.2	392.4
	15	40	NH-CH ₂ -	√√F	Н	350.8	350.4

Table 1. cont.

	5	#	Y	R ¹	R ²	М+Н са	lc'd
				\sim			
ja ra		41	NH-CH ₂ -	\sim	Н	389.2	388.5
		42	NH-CH ₂ -	√ Ç _F	н	351.0	350.4
H CO CO		43	NH-CH ₂ -	\sim \sim \sim	Н	367.1	366.8
ρ= ± /٦.		44	NH-CH ₂ -	- F	Н	401.3	400.4
Č N	10	45	NH-CH ₂ -		Н	377.2	376.5
		46	NH-CH ₂ -		Н	361.4	360.4
		47	NH-CH ₂ -		Н	377.1	376.4
		48	NH-CH ₂ -		Н	347.1	346.4
		49	NH-CH ₂ -	ОН	Н	349.1	348.4
	15	50	NH-CH ₂ -		Н	393.2	392.4

Table 1. cont.

5 M+H calc'd 51 NH-CH₂-411.2 411.3 403.1 401.3 52 NH-CH₂-53 NH-CH2-415.2 414.4 10 54 NH-CH₂-Н 393.2 392.4 55 NH-CH₂-403.2 401.3 56 NH-CH₂-Н 351.0 350.4 57 Н 369.1 366.8 NH-CH2-58 NH-CH-412.3 411.5 15 59 Н 338.8 338.4 NH-CH₂-

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Table 1. cont.

	5	#	Y	R ¹	R ²	М+Н са	lc'd
ļ.a		60	NH-CH ₂ -		Н	334.1	333.4
		61	NH-CH ₂ -		Н	333.6	333.4
(n (d 14		62	NH-CH ₂ -		Н	333.6	333.4
		63	NH-CH ₂ -	но	Н	361.1	360.4
	10	64	NH-CH ₂ -		н	379.0	378.4
		65	NH-CH ₂ -		Н	399	398.9
		66	NH-CH ₂ -		Ĭ	н 522.	3 521

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Example 67

5 {5-Fluoro-2-[(4-pyridylmethyl)amino](3-pyridyl)}-N-[4-(isopropyl)phenyl]carboxamide

[6-Chloro-5-fluoro-2-[(4-pyridylmethyl)amino](3pyridyl)}-N-[4-(methylethyl)phenyl]carboxamide (50 mg, 0.125 10 mmol, from Example 66) dissolved in EtOH (10 mL) with TEA (0.5 mL) and suspended with Pd/C (10%, 5 mg). The mixture was stirred at RT under a H2 balloon for 45 min. The mixture was filtered through a layer of Celite® and the filtrate was concentrated in vacuo. The residue was partitioned between CH2Cl2 and aq. NaHCO3 (sat.). The organic solution was dried over Na2SO4 and concentrated in vacuo to give the title compound. MS: 365 (M+1). Calc'd. for C21H21FN4O - 364.42.

Example 68

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2-[(Pyridin-4-ylmethyl)amino]-N-[4-tert-butyl-3-(1,2,3,6tetrahydropyridin-4-yl)phenyl](3-pyridyl)carboxamide

Step A Preparation of 2-bromo-1-tert-butyl-4-nitrobenzene

NBS (125.0 g, 697.5 mmol) was slowly added to a solution of TFA: H_2SO_4 (5:1, 750 mL) and tert-buty1-4-nitrobenzene (100.0 g, 558.0 mmol) at RT. The solution was stirred for 24 h then poured over 5 kg of ice. The resulting suspension was filtered and washed with a 1:1 MeOH: H_2O solution (200 mL) and dried in a vacuum oven. MS (ES+): 258.1, 260.1 (M+H) $^{\circ}$. Calc'd for $C_{10}H_{12}BENO_2$: 257.01.

Step B Preparation of 4-(2-tert-butyl-5-nitrophenyl)pyridine

To a solution of 2-bromo-1-tert-butyl-4-nitrobenzene (8.6 g, 33.3 mmol) and toluene (70 mL) in a 150 mL round bottom flask, 4-pyridylboronic acid (4.5 g, 36.6 mmol), Pd(PPh₃) $_4$ (3.8 g, 3.3 mmol) and K₂CO₃ (13.8 g, 99.9 mmol) were added. The solution was stirred for 24 h at 80°C before cooling to RT. The solution was filtered through a pad of Celite $^{\circ}$ and purified by silica flash chromatography (30% EtOAc/Hexanes). This afforded the desired compound as a yellow solid. MS (ES+): 257.2 (M+H) $^{\circ}$; (ES-): 255.2 (M-H) $^{\circ}$.

Calc'd for C15H16N2O2: 256.12.

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Step C Preparation of 4-(2-tert-butyl-5-nitrophenyl)-1-methylpyridinium

4-(2-tert-Butyl-5-nitrophenyl)pyridine (2.0 g, 7.8 mmol, Step B) was added to a round-bottom flask and dissolved in EtOH (10 mL). MeI (30 mL) was added and the flask was placed in an 80°C sand bath and heated to reflux. After 6 h the solution was cooled to RT and the excess MeI and EtOH was concentrated in vacuo resulting in the desired compound as a light brown solid. MS (ES+): 271.2 (M+H)'; (ES-): 269.2 (M-H)'. Calc'd for C₁₆H₁₀N₂O': 271.14.

Step D Preparation of 4-tert-butyl-3-(1-methyl-1,2,3,6-tetrahydropyridin-4-yl)aniline

4-(2-tert-Butyl-5-nitrophenyl)-1-methylpyridinium (2.1 g, 7.8 mmol, Step C) was added to a 100 mL round-bottom flask and dissolved in a 10% H2O/EtOH mixture. To the flask iron dust (1.31 g, 23.4 mmol) and NH₄Cl (460 mg, 8.6 mmol) were added. The flask was placed in a 100°C sand bath and heated to reflux. After 2 h the solution was cooled to RT and filtered through a pad of Celite. The resulting solution was concentrated in vacuo to a vellow solid and redissolved in MeOH (20 mL, anhydrous). The solution was cooled to 0°C by placing it in an ice bath and slowly adding NaBH4 (450 mg, 11.7 mmol). After addition of the NaBH4, the solution was cooled to RT and stirred for 30 min. solvent was concentrated in vacuo and the solid was redissolved in CH2Cl2 and filtered. The solution was again concentrated in vacuo to afford an amorphous clear yellow solid. MS (ES+): 245.2 (M+H)*. Calc'd for C16H24N2: 244.19.

Step E Preparation of 2-[(pyridin-4-ylmethyl)amino]-N-[4-tert-butyl-3-(1,2,3,6-tetrahydropyridin-4-yl)phenyl](3-pyridyl)carboxamide

3.0

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The titled compound was prepared from 4-tert-butyl-3-(1-methyl-1,2,3,6-tetrahydropyridin-4-yl) aniline (Step D) by the method described in Example 25. MS: (ES+) 456.3 (M+H); (ES-) 454.4 (M-H). Calc'd for $C_{26}H_{31}N_5O$ - 455.59.

Example 69

N-(3,4-Dichlorophenyl) {6-[(2-morpholin-4-ylethyl)amino]-2-[(4-pyridylmethyl)amino](3-pyridyl)}carboxamide

A mixture of N-(3,4-dichlorophenyl)(6-chloro-2-[(4-pyridylmethyl)amino](3-pyridyl))carboxamide (18 mg, 0.044 mmol, made from 2,6-dichloronicotinic acid) and 2-morpholin-4-ylethylamine (300 µL) was stirred at 80°C for 20 h. The reaction mixture was purified on silica gel chromatography to yield N-(3,4-dichlorophenyl)(6-[(2-morpholin-4-ylethyl)amino]-2-[(4-pyridylmethyl)-amino](3-pyridyl))carboxamide. MS (ES+): 501 (M+H)*; (ES-): 499 (M-

20 pyridyl))carboxamide. MS (ES+): 501 (M+H)*; (ES-): 499 (M-H) $^{-}$. Calc'd for $C_{24}H_{26}Cl_{2}N_{6}O_{2}$ - 500.15.

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Example 70

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N-[4-(Morpholin-4-ylmethyl)phenyl] {2-[(4pyridylmethyl)amino](3-pyridyl)}carboxamide

Step A Preparation of 4-[(4-nitrophenyl)methyl]morpholine

A mixture of nitrobenzyl bromide (648 mg, 3.0 mmol) and morpholine (522 mg, 6.0 mmol) in CH_2Cl_2 was stirred for 5 h at RT. Filtration to remove the white solid, and the filtrate was concentrated to give 4-[(4-nitrophenyl)-methyl]morpholine as a solid, which was used in next step without further purification.

Step B Preparation of 4-(morpholin-4-ylmethyl)phenylamine

A mixture of 4-[(4-nitrophenyl)methyl]morpholine (220 mg, 1.0 mmol, Step A), iron powder (279 mg, 5.0 mmol) and NH₂Cl (39 mg, 0.7 mmol) in EtOH (3 mL) and H₂O (3 mL) was stirred for 4 h at 80°C. Filtration and concentration gave the crude <math>4-(morpholin-4-ylmethyl)-phenylamine, which was used in next step without further purification.

25 Step C Preparation of N-[4-(morpholin-4-ylmethyl)phenyl](2-[(4-pyridylmethyl)amino](3-pyridyl))carboxamide

The titled compound was prepared from 4-(morpholin-4-ylmethyl)phenylamine (Step B) by the method described in

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Example 25. MS (ES+): 404 (M+H); (ES-): 402 (M-H). Calc'd. for $C_{22}H_{24}N_{4}O_{2}$ - 403.20.

Example 71

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N-(4-{2-[(tert-Butoxy)carbonylamino]ethyl}phenyl)(2-[(4-pyridylmethyl)amino](3-pyridyl))carboxamide

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Step A Preparation of (tert-butoxy)-N-[2-(4-nitrophenyl)ethyl]carboxamide

A mixture of 2-(4-nitrophenyl)ethylamine (1.01 g, 5.0 mmol), and di-tert-butyl dicarbonate (1.09 g, 5.0 mmol) in CH₂Cl₂ (20 mL) and 1N NaOH (20 mL) was stirred for 20 h at RT. The mixture was extracted with CH₂Cl₂, washed with brine, and dried with MgSO₄. Filtration and concentration yielded (tert-butoxy)-N-[2-(4-nitrophenyl)ethyl]carboxamide, which was used in next step without further purification.

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Step B Preparation of N-[2-(4-aminophenyl)ethyl](tert-butoxy)carboxamide

A mixture of (tert-butoxy)-N-[2-(4-nitrophenyl)ethyl]-carboxamide (570 mg, 2.15 mmol, Step A), iron powder (602 mg, 10.75 mmol) and NH₄Cl (82 mg, 1.5 mmol) in EtOH (6 mL) and H₂O (6 ml) was stirred for 4 h at 80°C. Filtration and concentration gave the crude compound, which was used in next step without further purification.

Step C Preparation of N-[4-(morpholin-4-ylmethyl)phenyl]{2-[(4-pyridylmethyl)amino](3-pyridyl)}carboxamide

The titled compound was prepared from N-[2-(4aminophenyl)ethyl](tert-butoxy)carboxamide (Step B) by the method described in Example 25. MS (ES+): 448 (M+H): (ES-): 446 (M-H). Calc'd. for C25H29N5O3 - 447.23.

Example 72

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N-[4-(2-Aminoethyl)phenyl] {2-[(4-pyridylmethyl)amino](3pyridyl) } carboxamide

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To the solution of N-(4-{2-[(tertbutoxy)carbonvlaminol-ethyl}phenyl){2-[(4pyridylmethyl)aminol(3-pyridyl))carboxamide (96 mg, 0.22 mmol, Example 71) in CH2Cl2 (3 mL) was added TFA (3 mL). The mixture was stirred for 3 h at RT. The reaction mixture was concentrated and dried in vacuo to yield N-[4-(2aminoethyl)phenyl]{2-[(4-pyridylmethyl)-amino](3pvridvl) carboxamide. MS (ES+): 348 (M+H); (ES-): 346 (M-H). Calc'd. for C20H21N5O - 347.17.

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The following compounds (Example a-m) were synthesized by the method described above, unless specifically described.

a) N-[3-(azetidin-3-vlmethoxy)-5-trifluoromethyl-phenyl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide.

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- b) 2-[(2,3-Dihydro-benzofuran-5-ylmethyl)-amino]-N-[3,3-dimethyl-1-(piperidin-4-ylmethyl)-2,3-dihydro-1H-indol-6-yl]-nicotinamide. M+H 512.3; Calc'd 511.7.
- c) N-[3-(piperazine-1-carbonyl)-5-trifluoromethyl-phenyl]-2-[(pyridin-4-vlmethyl)-amino]-nicotinamide. M+H 485.3.
- d) N-[3-(piperazine-1-methyl)-4-pentafluoroethyl-phenyl]-2-[(pvridin-4-vlmethyl)-amino]-nicotinamide. M+H 521.4.
- c) N-[3-(piperazine-1-methyl)-5-trifluoromethyl-phenyl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide. M+H 471.2; Calc'd 470.
- d) N-[1-(2-Amino-acetyl)-3,3-dimethyl-2,3-dihydro-1H-indol-6-yl]-2-[(2-methoxy-pyridin-4-ylmethyl)-amino)nicotinamide. M+H 461.1.
- e) N-[1-(2-Amino-acetyl)-3,3-dimethyl-2,3-dihydro-1H-indol-6-yl]-2-[(pyridin-4-ylmethyl)-amino]nicotinamide. M+H 431.4.
- f) (S) N-[3-(pyrrolidin-2-yl-methoxy)-4-pentafluoroethyl-phenyl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide. M+H 522.6; Calc'd 521.5.
- 20 g) (R) N-[3-(pyrrolidin-2-yl-methoxy)-4-trifluoromethyl-phenyl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide.
 M+H 472.6; Calc'd 471.5.
 - h) (R) N-[3-(pyrrolidin-2-yl-methoxy)-4-pentafluoroethyl-phenyl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide. M+H 522.3; Calc'd 521.5.
 - (S) N-[3-(pyrrolidin-2-yl-methoxy)-5-trifluoromethylphenyl]-2-[(pyridin-4-yl-methyl)-amino]-nicotinamide.
 M+H 472: Calc'd 471.5.
- j) (S) N-[3-(4-piperdinyloxy)-5-trifluoromethyl-phenyl]-230 [(pyridin-4-ylmethyl)-amino]-nicotinamide. M+H 472;
 Calc'd 471.5.
 - k) 2-[(2-Methoxy-pyridin-4-yl-methyl)-amino]-N-[3-(piperidin-4-yloxy)-5-trifluoromethyl-phenyl]nicotinamide.

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- N-{4-tert-Butyl-3-[2-(piperidin-4-yl)-methoxy]-phenyl}-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide. M+H 474.
- m) N-[4-tert-Butyl-3-(pyrrolidin-2-ylmethoxy)-phenyl]-2[(pyridin-4-ylmethyl)-amino]-nicotinamide. M+H 460.
- 5 n) 2-[(2-Methoxy-pyridin-4-ylmethyl)-amino]-N-[3-(pyrrolidin-2-ylmethoxy)-5-trifluoromethyl-phenyl]nicotinamide.
 - N-(3,3-Dimethyl-1-pyrrolidin-2-ylmethyl-2,3-dihydro-1H-indol-6-yl)-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide.

Example 73

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N-[4-(tert-Butyl)-3-nitrophenyl]{2-[(2-pyridylmethyl)amino](3-pyridyl)}carboxamide

MS (ES+): 406 (M+H); (ES-): 405 (M-H). Calc'd. for $C_{22}H_{23}N_5O_3$ 20 - 405.18.

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Example 74

N-[3-Amino-4-(tert-butyl)phenyl]{2-[(2-pyridylmethyl)amino](3-pyridyl)}carboxamide

A mixture of N-[4-(tert-buty1)-3-nitropheny1](2-[(210 pyridylmethy1)amino](3-pyridyl))carboxamide (100 mg, 0.25
mmol, Example 73), iron powder (69 mg, 1.25 mmol) and NH₄Cl
(10 mg, 0.17 mmol) in EtOH (0.5 mL) and H₂O (0.5 ml) was
stirred for 4 h at 80°C. The reaction mixture was filtered,
concentrated, and purified through column chromatography to
15 give the product. MS (ES+):376 M+H);(ES-):374(M-H). Calc'd.
for C₂₂H₃₅N₅O-375.21.

Example 75

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N-[4-(Isopropy1)pheny1]{2-[(2-pyridylmethy1)amino](3pyridy1)}carboxamide MS (ES+): 347 (M+H); (ES-): 345 (M-H). Calc'd. for $C_{21}H_{22}N_4O$ - 346.18.

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Example 76

N-(3-Aminosulfonyl-4-chlorophenyl) {2-[(4-pyridylmethyl)amino](3-pyridyl)}carboxamide

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MS (ES+): 418 (M+H); (ES-): 416 (M-H). Calc'd. for $C_{18}H_{16}N_5O_3S$ - 417.07.

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Example 77

 $\label{eq:N-substitution} $N-\{3-[(4-\text{methylpiperazinyl})sulfonyl]phenyl\} $\{2-[(4-\text{pyridylmethyl})amino](3-\text{pyridyl})\}$ carboxamide$

Step A Preparation of 3-[(4-methylpiperazinyl) sulfonyl]-1-nitrobenzene

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A mixture of 3-nitrobezenesulfonyl chloride (664 mg, 3.0 mmol) and methylpiperazine (600 mg, 6.0 mmol) in EtOH was stirred for 2 h at RT. The reaction was concentrated and triturated in Et₂O to yield a yellowish solid, 3-[(4-methylpiperazinyl) sulfonyl]-1-nitrobenzene, and was used in next step without further purification.

Step B Preparation of 3-[(4-

methylpiperazinyl) sulfonyl]phenylamine

3-[(4-Methylpiperazinyl)] sulfonyl]phenylamine was analogously synthesized from 3-[(4-methylpiperazinyl)] sulfonyl]-1-nitrobenzene (Step A) by the method described in Example 74, which was used in next step without further purification. MS (ES+): 256 (M+H). Calc'd. for $C_{11}H_17N_3O_2S-255.10$.

Step C Preparation of N-[4-(morpholin-4-ylmethyl)phenyl](2-[(4-pyridylmethyl)amino](3-pyridyl))carboxamide

The titled compound was prepared from 3-[(4-20 methylpiperazinyl)sulfonyl]phenylamine (Step B) by the method described in Example 25. MS (ES+): 467 (M+H); (ES-): 465 (M-H). Calc'd. for C₂₁H₂₆N₆O₁S - 466.18.

Example 78

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N-[4-(1,1,2,2,2-Pentafluoroethyl)phenyl]{2-[(4pyridylmethyl)amino](3-pyridyl)}carboxamide

Step A Preparation of 4-(1,1,2,2,2-

5 pentafluoroethyl)phenylamine

1-Nitro-4-(1,1,2,2,2,2-pentafluoroethyl) benzene was synthesized by the method described in the reference [John N. Freskos, Synthetic Communications, 18(9), 965-972 (1988)]. It was reduced with Fe similar to that described in Example 74. It was used in next step without further

Step B Preparation of N-[4-(1,1,2,2,2-

Pentafluoroethyl)phenyl]{2-[(4-pyridylmethyl)amino](3-

15 pyridyl)}carboxamide

purification.

The titled compound was prepared from 4-(1,1,2,2,2-pentafluoroethyl) phenylamine (Step A) by the method described in Example 25. MS (ES+): 423 (M+H); (ES-): 421 (M-H). Calc'd. for $C_{20H15}FN_{6}O$ - 422.12.

Example 79

N-[4-(1,1,2,2,3,3,4,4,4-Nonafluorobuty1)pheny1]{2-[(4-pyridylmethyl)amino](3-pyridyl)}carboxamide

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Step A Preparation of 4-(1,1,2,2,3,3,4,4,4-

nonafluorobutyl) phenylamine

The title intermediate was analogously synthesized by the method described of W. A. Gregory, et al. [J. Med.

Chem., 1990, 33, 2569-2578]. 1-nitro-4-(1,1,2,2,3,3,4,4-monofluorobutyl) benzene was reduced with Fe described in Example 68, Step D, and used in next step without further purification.

10 Step B Preparation of N-[4-(1,1,2,2,3,3,4,4,4nonafluorobutyl)phenyl](2-[(4-pyridylmethyl)amino](3pyridyl))carboxamide

The titled compound was prepared from 4-(1,1,2,2,3,3,4,4,4,4-nonafluorobutyl)phenylamine (Step A) by the method described in Example 25. MS (ES+): 523 (M+H); (ES-): 521 (M-H). Calc'd. for C₂₂H₁₅F₄N₄O-522.37.

Example 80

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N-[4-(Isopropyl)phenyl]{2-[(2-(1,2,4triazolyl)ethyl)amino](3-pyridyl)}carboxamide

25 Step A Preparation of 2-(1,2,4-triazolyl)ethylamine

A mixture of (tert-butoxy)-N-(2-chloroethyl)-carboxamide (900 mg, 5 mmol), 1,2,4-triazole (690 mg, 10

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mmol) and $Na_{3}CO_{3}$ (1.06 g, 10 mmol) in DMF (3 mL) was stirred overnight at $100^{\circ}C$. The mixture was filtered and concentrated to give an oil. The oil was treated with TFA (10 mL) and stirred for 3 h. The reaction was concentrated to give the titled intermediate, which was used in next step without further purification.

Step B Preparation of N-[4-(methylethyl)phenyl]{2-[(2(1,2,4-triazolyl)ethyl)amino](3-pyridyl)}carboxamide

The titled compound was prepared from 2-(1,2,4-triazolyl)ethylamine (Step A) by the method described in Example 25. MS (ES+): 351 (M+H); (ES-): 349 (M-H). Calc'd. for $C_{16}H_{27}N_6O$ - 350.19.

Example 81

(2-{[2-(2-Pyridylamino)ethyl]amino}(3-pyridyl))-N-[3-(trifluoromethyl)phenyl]carboxamide

Step A Preparation of 2-(2-pyridylamino)ethylamine

Ethylenediamine (6 g, 0.1 mol) and 2-fluoropyridine (10 g, 0.1 mol) were heated neat at 120° C overnight. The reaction was cooled and the residue was used in next step without further purification.

Step B Preparation of (2-{[2-(2-pyridylamino)ethyl]amino}-(3-pyridyl))-N-[3-(trifluoromethyl)phenyl]carboxamide

The titled compound was prepared from 2-(2-pyridylamino)ethylamine (Step A) by the method described in Example 25. MS (ES+): 402 (M+H); (ES-): 400 (M-H). Calc'd. for $C_{20}H_{18}F_{1}N_{2}O$ - 401.15.

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2-[(Pyridin-4-ylmethyl)-amino]-N-(3-trifluoromethyl-phenyl)nicotinamide

Step A: Preparation of (2-chloro(3-pyridyl))-N-(3-trifluoromethylphenyl)carboxamide

2-Chloropyridine-3-carbonyl chloride (18.02 g, 0.102 mol) in CH_2Cl_2 (100 ml) was added dropwise (via an addition funnel) to a stirred solution of 3-(trifluoromethyl)-aniline (15.00 g, 0.093 mol) and DIEA (24.39 ml, 0.14 mol) in CH_2Cl_2 (500 ml) at 0°C. The mixture gradually was warmed to RT. The reaction continued for 18 h before washing several times with saturated NaHCO₃ aqueous solution and brine, respectively. The organic layer was dried over Na_2SO_4 and evaporated. The resulting oil was purified over silica gel with EtOAc/hexane (2:1) as eluant to leave the amide as a white solid (26.08 g). MS: (ES+) 301 (M + 1)*; (ES-): 299 (M - 1)°. Calc'd for $C_{11}H_2Cl_5N_5O$: 300.03.

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Step B: Preparation of N-[3-trifluoromethylphenyl phenyl]{2-[(4-pyridylmethyl)amino](3-pyridyl)}carboxamide hydrochloride

The amide (10.0 g 0.033 mol, Step A) and 4aminomethylpyridine (10.81 g, 0.10 mol) were combined and heated at 120°C for 4 h. After cooling to RT, the residue was dissolved in EtOAc and washed several times with saturated NaHCO, aqueous solution and brine, respectively. The organic layer was dried over Na2SO4 and evaporated. The 10 crude vellow oil was purified over silica gel with EtOAc as eluant to leave an amber oil (10.9 g). The free base was dissolved in MeOH (20 ml) and treated with a HCl ethereal solution (1.0 eq.). The solvent was evaporated to leave the salt as a white solid. The HCl salt was dried in vacuo at 30° C for 24 h. MS: (ES+) 373 (M + 1)⁺; (ES-): 371 (M - 1)⁻.

15 Calc'd. for $C_{19}H_{15}F_3N_4O - 372.12$.

The following compounds (Examples 83-138) were analogously synthesized by the method described in Example 82 unless specifically described. Detailed intermediate preparations are included.

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Table 2.

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#	Y	R ¹	R ²	M+H	calc'd
83	-NH-CH ₂ -	h	Н	356	355.14
84	-NH-CH ₂ -		н	431	430.12
85	-NH-CH₂-		н	359	358.1
86	-NH-CH ₂ -		Н	355	354.15
87	-NH-CH ₂ -		н	359	358.18
88	-NH-CH ₂ -		Н	349	348.128
89	-NH-CH ₂ -		Н	355	354.15
90	-NH-CH ₂ -		н	395	394.18

Table 2 cont.

5	#	Y	R ¹	R ²	M+H	calc'd
	91	-NH-CH ₂ -	H ₃ C	Н	339	338.12
	92	-NH-CH ₂ -		Н	339	338.12
	93	-NH-CH ₂ -		Н	345	344.16
	94	-NH-CH ₂ -		Н	361	360.20
10	95	-NH-CH ₂ -	, CH _A	Н	361	360.20
	96	-NH-CH ₂ -	F F	Н	319	318.5
	97	-NH-CH ₂ -		Н	389	388.11
	98	-NH-CH ₂ -	СНэ	Н	333	332.16

Table 2. cont

5	#	Y	R ¹	R²	M+H	calc'd
	99	-NH-CH ₂ -	CH ₀	Н	361	360.2
	100	-NH-CH ₂ -		Н	431	430
	101	-NH-CH ₂ -		Н	349	348.16
	102	-NH-CH ₂ -	CH ₃	н	333	332.16
10	103	-NH-CH₂-		н	457	456.18
	104	-NH-CH ₂ -	Ü	н	381	380.16
	105	-NH-CH ₂ -		Н	395	394.18
	106	-NH-CH ₂ -	CH ₃	Н	334	333.16
15	107	-NH-CH ₂ -	СНа	Н	348	347.17

Table 2. cont.

5	#	Y	R ¹	R ²	M+H	calc'd
			H ₃ C CH ₃	٠		
	108	-NH-CH ₂ -		Н	362	361.19
	109	-NH-CH ₂ -	H ₃ C CH ₃	Н	321	320.13
	110	-NH-CH ₂ -	ÇF ₃	Н	348	347.17
	111	-NH-CH ₂ -	CF ₃	н	441	440.11
10	112	-NH-CH ₂ -	CF ₃	Н	407	406.08
	113	-NH-CH ₂ -		Н	353	352.11
	114	-NH-CH ₂ -		н	383	382.11

Table 2. cont.

5						
	#	Y	R ¹	R ²	M+H	calc'd
	115	-NH-CH ₂ -	H ₃ C	Н	388	387.43
	116	-NH-CH₂-	LT _N	н	346	345.00
	117	-NH-CH ₂ -		н	344	343.38
10	118	-NH-CH ₂ -	₩ NH	н	344	343.38
	119	-NH-CH ₂ -	HN-N H	н	344	343.38
	120	-NH-CH ₂ -		Н	351	350.43
	121	-NH-CH ₂ -	HN-N	Н	371	370.43

Example 122

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N-(4-Phenoxyphenyl) {2-[(2-(2-pyridyl)ethyl)amino](3pyridyl)}carboxamide

MS: 411 (M+1); 409 (M-1). Calc'd. for $C_{25}H_{22}N_4O_2$ - 410.17.

Example 123

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2-[(Benzo[b]thiophen-3-ylmethyl)amino](3-pyridyl))-N-(4phenoxyphenyl)carboxamide

MS: (ES+) 452 (M + 1)*; (ES-): 450 (M - 1)*. Calc'd. for $C_{27}H_{21}N_3O_2S$ - 451.14.

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Example 124

N-{2-[2-(Dimethylamino)ethoxy]-5-(tert-butyl)phenyl}{2-[(4pyridylmethyl)amino](3-pyridyl)}carboxamide

Step A - Preparation of (2-[4-(tert-buty1)-2-nitrophenoxy]ethyl)dimethylamine

To a mixture of 2-nitro-4-tert-butylphenol (2 g) and N,N-dimethylethanolamine (1.3 g) and PPh₃ (4 g) in THF (50 ml) was added DEAD (2.6 ml). The reaction was stirred at RT for 1 h, diluted with EtOAc (50 ml) and washed with 1 N HCl twice. The aqueous layer was basified with NaHCO₃, extracted with EtOAc twice and washed with H_2O and brine. The organic layer was dried over Na_2SO_4 and evaporated to give (2-[4-(tert-butyl)-2-nitrophenoxy]-ethyl)dimethylamine, was used in next step without further purification.

20 <u>Step B - Preparation of {2-[4-{tert-buty1}-2-aminophenoxy}-</u> ethyl}dimethylamine

(2-[4-(tert-Butyl)-2-nitrophenoxy]-ethyl) dimethylamine (Step A) was hydrogenated under H₂ atmosphere to give (2-[4-(tert-butyl)-2-aminophenoxy]-

25 ethyl}dimethylamine, and used in next step without further purification.

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Step C - Preparation of N-{2-[2-(dimethylamino)ethoxy]-5-(tert-butyl)phenyl}{2-[(4-pyridylmethyl)amino](3pyridyl) } carboxamide

The titled compound was prepared from {2-[4-(tertbutyl)-2-aminophenoxy]-ethyl}dimethylamine (Step B) by the method described in Example 82. MS (ES+): 448 (M+H); (ES-): 446 (M-H). Calc'd. for $C_{26}H_{33}N_5O_2$ - 447.26.

Example 125

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$N-[4-(tert-Buty1)-3-(4-methylpiperaziny1)pheny1]{2-[(4-methylpiperaziny1)pheny1]}$ pyridylmethyl)amino](3-pyridyl)}carboxamide

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Step A - Preparation of 1-[2-(tert-butylphenyl]-4methylpiperazine

A mixture of 2-tert-butylaniline (5.4 g) and Nmethylbis(2-chloroethyl)amine hydrochloride (7 g) and K_2CO_3 (5 g) in NaI (2 g) in diglyme (150 ml) was heated at 170°C for 8 h. The reaction was filtered and the filtrate was evaporated under high vacuum. The residue was mixed with EtOAc (200 ml) and ${\rm H}_2{\rm O}$ (200 ml) and extracted with EtOAc twice. The combined organic layer was washed with brine, 25 dried over Na₂SO₄ and evaporated to give crude 1-[2-(tertbutylphenyl]-4-methyl-piperazine, which was used in next step without further purification.

Step B - Preparation of 1-[2-(tert-butyl)-5-aminophenyl]-4-methylpiperazine

The crude 1-[2-(tert-butylphenyl]-4-methylpiperazine (260 mg, Step A) was stirred with H₂SO₄ (3 ml) at 0°C and HNO₃ (1.2 ml) was slowly added to the reaction. The reaction was warmed to RT, stirred for 30 min. and poured on ice and basified with K₂CO₃ slowly. The solution was extracted with EtOAc three times, washed with H₂O, followed by brine, dried over Na₂SO₄, and evaporated under reduced pressure. The residue was purified by column chromatography to give 1-[2-(tert-butyl)-5-nitrophenyl]-4-methylpiperazine (260 mg), which was hydrogenated under H₂ atmosphere to give 1-[2-(tert-butyl)-5-mitrophenyl]-4-methylpiperazine.

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Step C - Preparation of N-[4-(tert-Butyl)-3-(4methylpiperazinyl)phenyl){2-[(4-pyridylmethyl)amino](3pyridyl)}carboxamide

The titled compound was prepared from 1-{2-(tert-20 buty1)-5-aminopheny1}-4-methylpiperazine (Step B) by the method described in Example 82. MS (ES+): 459 (M+H); (ES-): 457 (M-H). Calc'd. for C₂₇H₃₄N₆O - 458.28.

Example 126

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N-[3-(4-Methylpiperazinyl)phenyl]{2-[(4-pyridylmethyl)amino](3-pyridyl)}carboxamide

5 Step A - Preparation of 3-(4-methylpiperazinyl)phenylamine

The intermediate was analogously synthesized from 3-nitroaniline by the method described in Example 130.

Step B - Preparation of N-[3-(4-methylpiperazinyl)phenyl]{2-

10 [(4-pyridylmethyl)amino](3-pyridyl)}carboxamide

The titled compound was prepared from 3-(4-methylpiperazinyl)phenylamine (Step A) by the method described in Example 82. MS (ES+): 403 (M+H); (ES-): 401 (M-H). Calc'd. for C₂₃H₂₆N₆O - 402.22.

Example 127

N-[4-(4-Methylpiperaziny1)phenyl]{2-[(4-pyridylmethyl)amino](3-pyridyl)}formamide

Step A - Preparation of 4-methyl-1-(4-nitrophenyl)piperazine 1-Fluoro-4-nitrobenzene (3.0 g. 0.021 mol) and 1-

methylpiperazine (6.98 ml, 0.63 mol) were combined and heated neat at 90°C for 48 h. Upon cooling to RT, the resulting brown oil solidified. The crude material was purified by re-crystallization from EtOAc/Hexane mixtures to leave the title compound as an orange solid (3.59 g). MS:

 (ES+) 222 (M + 1)*; (ES-): 220 (M - 1) $^{-}$. Calc'd for $C_{11}H_{15}N_3O_2$: 221.12.

Step B - Preparation of 4-methyl-1-(4-aminophenyl)piperazine

4-Methyl-1-(4-nitrophenyl)piperazine (2.0 g, 9 mmol, Step A) and 10% Pd/C (200 mg) were added to EtOH/MeOH (1:1) (50 ml) at RT. The reaction stirred under a $\rm H_2$ atmosphere (via balloon) overnight. The mixture was filtered through a plug of Celite[®] and the filtrate was concentrated under reduced pressure to leave the desired material as a light yellow oil. The material was used in subsequent reaction without purification. MS: (ES+) 192 (M + 1)*; (ES-): 190

15 Step C Preparation of N-[4-(4-methylpiperazinyl)phenyl](2-[(4-pyridylmethyl)amino](3-pyridyl))formamide

(M - 1)". Calc'd for C11H17N3: 191.14.

The titled compound was prepared from 4-methyl-1-(4-aminophenyl)piperazine (Step B) by the method described in Example 82. MS (ES+): 403 (M+H); (ES-): 401 (M-H). Calc'd. for $C_{21}H_{26}N_6O$ - 402.22.

Example 128

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N-[1-(1-Methyl-(4-piperidyl))indolin-6-yl]{2-[(4-pyridylmethyl)amino](3-pyridyl)}carboxamide

Step A - Preparation of 1-(1-methyl(4-piperidyl))-630 nitroindoline

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6-Nitroindoline (5 g) was dissolved in 200 mL of dichloroethane, N-methyl-4-piperidone (5 g) was added to the mixture, followed by 12 g NaBH(OAc), and 1 mL of glacial AcOH. The mixture was stirred at RT overnight. Saturated NaHCO3 solution (200 mL) was added to the reaction mixture and stirred for 1 h. The resulting mixture was separated by separation funnel, the organic layer was extracted once with saturated NaHCO3 solution and once with brine. The resulting organic layer was dried over MgSO4, filtered and

10 concentrated in vacuo. The crude material was purified by flash chromatography on silica gel with 2:1 EtOAc:MeOH to afford an orange oil. MS: 262 (M+1). Calc'd. for $C_{14}H_{19}N_3O_{2}$ -261 32

15 Step B - Preparation of 1-(1-methyl-4-piperidyl)indoline-6ylamine

25 Step C - Preparation of N-{1-(1-methyl(4-piperidyl))indolin-6-v1)(2-[(4-pvridylmethyl)amino)(3-pvridyl)}carboxamide

The titled compound was prepared from 1-(1-methyl-4-piperidyl)indoline-6-ylamine (Step B) by the method described in Example 82. MS: 443 (M+1). Calc'd. for $C_{26}H_{16}N_6O-442.56$.

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Example 129

N-[1-(1-Methyl-(4-piperidyl))indolin-6-yl]{2-[(2-(3-

MS: 457 (M+1). Calc'd. for C27H32N6O-456.58.

Example 130

pyridyl)ethyl)amino](3-pyridyl)}carboxamide

N-[1-(2-Piperidylethyl)indolin-6-yl]{2-[(4-pyridylmethyl)amino](3-pyridyl)}carboxamide

Step A - Preparation of 1-(6-nitroindoliny1)-2piperidylethan-1-one

6-Nitroindoline (2.5 g) was dissolved in 200 mL of CH_2Cl_2 , followed by DIEA (2.5 g). The mixture was cooled down to 0°C in ice bath. Chloroacetyl chloride (1.7 g) in

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20 mL CH_2Cl_2 was added dropwise to the mixture over 10 min and the mixture was stirred at RT overnight. The mixture was extracted once with saturated $NaHCO_3$ solution and once with brine, the resulting organic layer was dried over $MgSO_4$,

- brine, the resulting organic layer was dried over MgSO₄,

 5 filtered and concentrated in vacuo. The crude material was
 purified by flash chromatography on silica gel with 3:2

 Hexane:EtOAc to afford a yellow oil (1.4 g) which was added
 to piperidine (5 mL), followed by NaI (100 mg). The mixture
 was heated at 70°C overnight then concentrated in vacuo and

 10 extracted between EtOAc and saturated NaHCO₃ solution, the
 organic layer was washed with brine, the resulting organic
 layer was dried over MgSO₄, filtered and concentrated in
 vacuo. The crude material was purified by flash
- chromatography on silica gel with 9:1 EtOAc:MeOH to afford a 15 yellow oil. MS: 290 (M+1). Calc'd. for C15H19N101-289.33.

Step B - Preparation of 1-(2-piperidylethyl)indoline-6vlamine

- 1-(6-Nitroindolinyl)-2-piperidylethan-1-one (1.6 g,
- Step A) was dissolved in 100 mL MeOH, the mixture was bubbled with N₂ for 10 min. 10% Pd/C (200 mg) was added and the mixture was stirred under H₂ overnight. The mixture was filtered through Celite® and concentrated in vacuo to afford a yellow solid. 400 mg was dissolved in 20 mL anhydrous THF,
- 5 mL borane-THF (1 M) solution was added dropwise and the mixture was stirred at RT overnight. The mixture was quenched with MeOH, 100 mg NaOH added and heated at 70°C for 30 min. The resulting mixture was concentrated in vacuo and extracted between EtOAc and saturated NaHCO3 solution, the
- 30 organic layer was washed with brine, the resulting organic layer was dried over $MgSO_4$, filtered and concentrated in vacuo to afford a yellow oil. MS: 246 (M+1). Calc'd. for $C_{15}H_{23}N_3-246.36$.

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Step C - Preparation of N-[1-(2-piperidylethyl)indolin-6y1]{2-((4-pyridylmethyl)amino](3-pyridyl)}carboxamide

The titled compound was prepared from 1-(2-piperidylethyl)indoline-6-ylamine (Step B) by the method described in Example 82. MS: 457 (M+1). Calc'd. for $C_{27}H_{12}N_6O-456.58$.

Example 131

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N-[1-(2-Piperidylacetyl)indolin-6-yl]{2-[(4-pyridylmethyl)amino](3-pyridyl)}carboxamide

15 MS:

MS: 471 (M+1). Calc'd. for $C_{27}H_{30}N_6O_2-470.57$.

Example 132

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N-[3,3-Dimethyl-1-(1-methyl(piperid-4-yl)indolin-6-yl](2-[(4-pyridylmethyl)amino](3-pyridyl))carboxamide

Step A - Preparation of N-(2-bromo-5-nitrophenyl)acetamide

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2-Bromo-5-nitroaniline (10 g) was dissolved in 500 mL of CH₂Cl₂, DIEA (6.6 g) was added to the mixture, followed by DMAP (100 mg). The mixture was cooled to 0°C in ice bath. Acetyl chloride (4 g in 50 mL CH₂Cl₂) was added dropwise to the reaction mixture. After the mixture was stirred at RT over 3 h, extracted once with saturated NaHCO₃ solution and once with brine, the resulting organic layer was dried over MgSO₄, filtered and concentrated in vacuo. The crude material was purified by flash chromatography on silica gel with 1:1 EtOAc:Hexane to 100% EtOAc to afford N-(2-bromo-5-nitrophenyl)acetamide as a white solid. MS: 258 (M-1). Calc'd. for CaHaBrNO₁-259.06.

Step B - Preparation of N-(2-bromo-5-nitrophenyl)-N-(2-methylprop-2-enyl)acetamide

A suspension of 2 g NaH (95% powder) in anhydrous DMF (100 mL) was cooled to -78° C, N-(2-bromo-5-nitrophenyl) acetamide (7 g, Step A) in dry DMF (50 mL) was added to the mixture under N₂ atmosphere. After the mixture was warmed to 0°C, 3-bromo-2-methylpropene (7.3 g in 20 dry DMF) was added to the mixture. The mixture was stirred at RT overnight. Next morning, the mixture was poured into a container of ice and extracted between saturated NaHCO₃ solution and EtOAc. The resulting organic layer was dried over MgSO₄, filtered and concentrated in vacuo. The crude material was purified by flash chromatography on silica gel with 7:2 hexane:EtOAc to afford the title compound as a yellow gum. MS: 314 (M+1). Calc'd. for C₁₂H₁₃BrN₂O₃-313.15.

30 Step C - Preparation of 1-(3,3-dimethyl-6-nitro-2,3-dihydro-indol-1-yl)ethanone

N-(2-Bromo-5-nitrophenyl)-N-(2-methylprop-2enyl)acetamide (4.5 g, Step B) was dissolved in anhydrous DMF (50 mL), tetraethyl-ammonium chloride (2.5 g), sodium

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formate (1.2 g), NaOAc (3 g) were added, and the resulting mixture was bubbled with N₂ gas for 10 min. Pd(OAc)₂ (350 mg) was added and the mixture was heated at 80°C under N₂ atmosphere overnight. After the mixture was concentrated in vacuo, it was partitioned between saturated NaHCO₃ solution and EtoAc, the resulting organic layer was dried over MgSO₄, filtered and concentrated in vacuo. The crude material was purified by flash chromatography on silica gel with 2:1 Hexane:EtoAc to afford the title compound as a yellow gum. MS: 235 (M+1). Calc'd. for CoH₁₀NO₂-234.25.

Step D - Preparation of 3,3-dimethyl-6-nitroindoline

1-(3,3-Dimethyl-6-nitro-2,3-dihydro-indol-1-yl)ethanone (1.8 g, Step C) was dissolved in EtOH (50 mL), 12N HCl (50 mL) was added and the resulting mixture was heated at 70°C overnight. After the mixture was concentrated in vacuo, it was partitioned between saturated NaHCO₃ solution and EtOAc, the resulting organic layer was dried over MgSO₄, filtered and concentrated in vacuo to afford a yellow solid. MS: 193 (M+1). Calc'd. for CuH1-NO-192.21.

Step E - Preparation of 3,3-dimethyl-1-(1-methyl-piperidin-4-yl)-6-nitro-2,3-dihydro-1H-indole

3,3-Dimethyl-6-nitroindoline (0.8 g) was dissolved in CH_2Cl_2 (50 mL), N-methyl-4-piperidone (1 g) was added to the mixture, followed by 2.5 g NaBH(OAc), and glacial AcOH (1 mL). The mixture was stirred at RT overnight. Saturated NaHCO, solution (50 ml) was added to the reaction mixture and stirred for 1 h. The resulting mixture was separated by separation funnel, the organic layer was extracted once with saturated NaHCO, solution and once with brine, the resulting organic layer was dried over MgSO₄, filtered and concentrated in vacuo. The crude material was purified by

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flash chromatography on silica gel with 9:1 EtOAc:MeOH to afford the title compound as an orange oil. MS: 290 (M+1). Calc'd. for ChillyNhOr-289.37.

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5 Step F - Preparation of 3,3-dimethyl-1-(1-methyl(4-piperidyl))indoline-6-vlamine

3,3-Dimethyl-1-(1-methyl-piperidin-4-yl)-6-nitro-2,3-dihydro-1H-indole (600 mg, Step E) was dissolved in MeOH (20 mL), the mixture was bubbled with $m H_2$ for 10 min. m 10%~Pd/C

10 (100 mg) was added and the mixture was stirred under H₂ overnight. The mixture was filtered through Celite® and concentrated in vacuo to afford the title compound as an oil. MS: 260 (M+1). Calc'd. for C₁₆H₂₅N₃-259.39.

The titled compound was prepared from 3,3-dimethyl-1-(1-methyl(4-piperidyl))indoline-6-ylamine (Step E) by the method described in Example 82. MS: 471 (M+1). Calc'd. for $C_{28}H_{14}N_{8}O-470.61$.

Example 133

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N-(3,3-Dimethylindolin-6-yl) {2-[(4-pyridylmethyl)amino](3pyridyl)}carboxamide

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Step A - Preparation of 1-acetyl-6-amino-3,3dimethylindoline

1-Acetyl-3.3-dimethyl-6-nitroindoline (250 mg) was dissolved in MeOH (20 mL), the mixture was bubbled with Ho for 10 min. 10% Pd/C (50 mg) was added and the mixture was stirred under H2 overnight. The mixture was filtered through Celite® and concentrated in vacuo. The crude material was purified by flash chromatography on silica gel with 1:1 EtOAc:CH2Cl2 to afford the title compound as a 10 white crystalline material. MS: 205 (M+1). Calc'd. for C12H16N2O-204.27.

Step B - Preparation of N-(1-acetyl- 3,3-dimethylindolin-6yl) {2-[(4-pyridylmethyl)amino](3-pyridyl)}carboxamide

The titled compound was prepared from 1-acetyl-6amino-3,3-dimethylindoline (Step A) by the method described in Example 82.

Step C - Preparation of N-(3,3-dimethylindolin-6-yl) {2-[(4pyridylmethyl)amino](3-pyridyl)}carboxamide

The titled compound was prepared from N-(1-acetyl-3,3-dimethylindolin-6-yl) {2-[(4-pyridylmethyl)amino](3pyridyl) carboxamide (Step B) by the deacylation method described in Example 993. MS: 374 (M+1). Calc'd. for C22H23N5O-373.45.

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Example 134

N-[3-(1-Methyl-(4-piperidyl))indol-5-yl]{2-[(4-pyridyl)methyl)aminol(3-pyridyl)}carboxamide

Step A - Preparation of 3-(1-methyl-1,2,3,6-tetrahydropyridin-4-yl)-5-nitro-1H-indole

5-Nitroindole (2.6 g) was dissolved in anhydrous MeOH (100 ml), followed by N-methyl-4-piperidone (5 g) and NaOMe powder (5 g). The mixture was heated to reflux under N₂ overnight. The mixture was concentrated in vacuo. The crude was partitioned between saturated NaHCO₃ solution and EtOAc, the resulting organic layer was dried over MgSO₄, filtered and concentrated in vacuo to afford a yellow solid. This solid was washed with EtOAc (5 mL) and MeOH (2 ml) to afford the title compound as a bright yellow solid. MS: 258 (M+1). Calc'd. for CuH₃NO₂-257.29.

Step B - Preparation of 3-(1-methyl-4-piperidyl)indole-5ylamine

3-(1-Methyl-1,2,3,6-tetrahydro-pyridin-4-yl)-5-nitro25 1H-indole (2.7 g, Step A) was dissolved in MeOH (50 mL), the mixture was bubbled with H₂ for 10 min. 10% Pd/C (150 mg) was added and the mixture was stirred under H₂ overnight.

The mixture was filtered through Celite® and concentrated in

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vacuo to afford 3-(1-methyl-4-piperidyl)indole-5-ylamine as a vellow oil. MS: 230 (M+1). Calc'd. for C12H10N1-229.32.

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Step C - Preparation of N-[3-(1-methyl-(4-piperidyl))indol-5-v1](2-((4-pyridylmethyl)amino)(3-pyridyl))carboxamide

The titled compound was prepared from 3-(1-methyl-4piperidvl)indole-5-vlamine (Step B) by the method described in Example 82. MS: 441 (M+1). Calc'd. for $C_{26}H_{28}N_6O-440.54$.

Example 135

N-[4-(1,1-Dimethyl-3-morpholin-4-vlpropyl)phenyl]{2-[(4pyridylmethyl)amino](3-pyridyl)}carboxamide

Step A - Preparation of methyl 2-methyl-2-(4nitrophenyl)propionate

To a stirred solution of 2-(4-nitrophenyl)-propionic acid (9 g, 46 mmol) in MeOH (300 mL) was added HCl (4M in dioxane, 11.5 mL, 46 mmol). The mixture was stirred at RT overnight and guenched with agueous NaHCO1. The mixture was extracted with EtOAc. The organic layer was dried over MgSO. evaporated under reduced pressure and to the partial residue at 0°C in THF (100 mL) was added NaH (1.66 g, 41.5 mmol). The mixture was stirred at RT for 1 h and MeI (2.58 g, 41.5 mmol) was added. The reaction was stirred at RT overnight and was quenched with H2O. The mixture was extracted with EtOAc, the organic layer was dried over MgSO4, evaporated under reduced pressure to give the title

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compound which was used in the next step without further purification. Calc'd for C11H13NO4: 223.08.

Step B - Preparation of 2-methyl-2-(4-nitro-phenyl)-propan1-o1

To a stirred solution of methyl 2-methyl-2-(4-nitrophenyl)propionate (5.32 g, 23.8 mmol, Step A) in THF (200 mL) at 0°C was added a solution of BH, 1M in THF (25.8 mL, 45.8 mmol). The reaction was stirred at RT overnight and quenched with MeOH. THF was evaporated under reduced pressure and the residue was diluted in EtOAc and aqueous 1M HCl was added. The mixture was extracted with EtOAc, the organic layer was dried over MgSO4 and evaporated under reduced pressure. The product was purified by flash chromatography using 40% EtOAc-hexane to give the title compound as a yellow solid.

Step C - Preparation of 2-methyl-2-(4-nitro-phenyl)propionaldehyde

To a stirred solution of the alcohol (2.08 g, 10.8 mmol, Step B) at 0°C in CH_2Cl_2 was added NMO (1.9 g, 16.1 mmol), molecular sieves 4Å and TPAP (76 mg, 0.2 mmol). The reaction was stirred for 1 h and was filtered on silica pad. Solvent was evaporated under reduced pressure. Crude aldehyde was used without further purification in the next step.

Step D - Preparation of 3-methyl-3-(4-nitrophenyl)butan-1-aldehyde

To a suspension of methoxymethyltriphenyl-phosphonium chloride (6.4 g, 18.6 mmol) in THF (150 mL) was added a solution of KHMDS 0.5 M in toluene (37 mL, 18.5 mmol). The mixture was stirred for 30 min and crude aldehyde (Step C) was added. The reaction was stirred at RT for 1 h and

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quenched with $\rm H_{2}O$. Mixture was extracted with EtOAc, dried and evaporated under reduced pressure. Et₂O was added and the formed precipitate was filtered on silica pad (rinsed with 40% EtOAc-hexane). The solvent was removed and crude product was dissolved in $\rm CH_{2}Cl_{2}$. A solution of TFA- $\rm H_{2}O$ (1:1, 10 mL) was added and the reaction was stirred for 2 h at RT. Aqueous NaHCO₃ was added until pH 7 and residue was extracted with $\rm CH_{2}Cl_{2}$. Organic layer was dried, filtered and evaporated. Crude compound was purified by flash

10 chromatography (40% EtOAc-hexane) to give the title compound as a yellow oil. Calc'd for C₁₁H₁₃NO₃: 207.09.

Step E - Preparation of 4-[3-methyl-3-(4-nitro-phenyl)butyl]-morpholine

To a stirred solution of 3-methyl-3-(4-nitrophenyl)butan-1-aldehyde (509 mg, 2.4 mmol, Step D) and morpholine (0.21 mL, 2.4 mmol) in THF (30 mL) was added NaBH(OAc)₃ (0.73 g, 3.4 mmol). The mixture was stirred at RT overnight and was washed with 1M HCl. CH₂Cl₂ was added and the layers were separated. The aqueous layer was basified to pH 9 using 1M NaOH and extracted with CH₂Cl₂. This organic layer was dried and evaporated yielding the morpholino compound. Calc'd for Cl₂H₂N₂O₃: 278.16.

25 Step F Preparation of 4-(1,1-dimethyl-3-morpholin-4ylpropyl)phenylamine

To a solution of 4-[3-methyl-3-(4-nitro-phenyl)-butyl]-morpholine (0.50g, 1.8 mmol, Step E) in THF (40 mL) was added AcOH (1.97 mmol, 34.5 mmol) followed by zinc (9.1 g, 137 mmol). The mixture was stirred for 1 h and filtered on Celite. The mixture was diluted with H_2O , and aqueous NaHCO₃ and the THF was evaporated. The residue was extracted with EtOAc, dried and evaporated to give the title intermediate. Calc'd for $C_{15}H_{23}N_{2}O$: 248.19.

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Step G - Preparation of N-[4-(1,1-dimethyl-3-morpholin-4-ylpropyl)phenyl](2-[(4-pyridylmethyl)amino](3-pyridyl))carboxamide

The titled compound was prepared from 4-(1,1-dimethyl-3-morpholin-4-ylpropyl)phenylamine (Step F) by the method described in Example 82. MS: 460.0 (M+1). Calc'd. for C₂-H₁N₁O₂ - 459.60.

Example 136

N-[4-(tert-Butyl)phenyl]{2-[({2-[(1-methyl(4-piperidyl))-methoxy](4-pyridyl)}methyl)amino](3-pyridyl)}carboxamide

Step A - Preparation of 4-hydroxymethyl-1-methylpiperidine

To a solution of 4-piperidylmethanol (1.0 g, 8.7 mmol) and HCHO (2 mL, 25 mmol, 37% in $\rm H_2O$) in CH₃CN was added NaCNBH₃ (0.5 g, 12.5 mmol). The resulting mixture was stirred for 1 h and filtered. The filtrate was concentrated and the residue was distilled (105°C, 40 torr) to give the title intermediate.

25 Step B - Preparation of {2-[(1-methyl-4-piperidyl)methoxy]-4-pyridyl}methylamine

To a suspension of NaH (0.44 g, 12.7 mmol, 60% in mineral oil) in DMF (25 mL) was added a solution of alcohol

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(1.1 g, 8.5 mmol, Step A) in 3 mL of DMF. After 20 min, a solution of 2-chloro-4-cyanopyridine (1.2 g, 8.5 mmol) in 2 mL of DMF was added. The resulting mixture was stirred for 2 h, diluted with CH₂Cl₂, and washed with H₂O twice. The organic layer was dried over Na₂SO₄ and concentrated to give 2-[(1-methyl-4-piperidyl)methoxy]pyridine-4-carbonitrile, which was hydrogenated under regular conditions to furnish the title intermediate. MS (ES+): 236 (M+H)⁴. Calc'd C₁₁H₂N₃O- 235.33.

Step C - Preparation of N-[4-(tert-butyl)phenyl]{2-[(1-methyl(4-piperidyl))-methoxy](4-pyridyl)}methyl)amino](3-pyridyl)}carboxamide

The title compound was prepared from $\{2-[(1-\text{methyl-4-piperidyl})\text{methoxy}]-4-pyridyl)\text{methylamine (Step B) by the method described in Example 82. MS (ES+): 488 (M+H)*; (ES-): 486 (M-H)^-. Calc'd <math>C_{29}H_{37}N_{5}O_{2}$ - 487.64.

Example 137

N-(4-Bromo-2-fluorophenyl) {2-[(4-pyridylmethyl)amino](3-pyridyl)}carboxamide

MS (ES+): 402 (M+H) $^{+}$; (ES-): 400. Calc'd C₁₈H₁₄BrFN₄O-401.238.

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Example 138

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N-[4-(tert-Butyl)phenyl](2-{[(2-chloro(4-pyridyl))methyl]amino}(3-pyridyl))carboxamide

MS (ES+): 395 (M+H) $^+$; (ES-): 393(M-H) $^-$. Calc'd $C_{22}H_{23}ClN_4O-10$ 394.90.

Example 139

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{2-[({2-[3-(Dimethylamino)prop-1-ynyl](4pyridyl))methyl)amino](3-pyridyl))-N-[4-(tertbutyl)phenyl]carboxamide

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A mixture of N-[4-(tert-butyl)phenyl](2-{((2-chloro(4-pyridyl))methyl]amino)(3-pyridyl))carboxamide (0.15 g, 0.38 mmol, Example 139), 1-dimethylamino-2-propyne (62 mg, 0.76 mmol), PdCl₂(PPh₃)₂ (13 mg, 0.0019 mmol) and CuI (7 mg, 0.019 mmol) in 1 mL of TEA was heated at 100°C in a sealed tube for 3 h. The resulting mixture was filtered over Celite[©]. The filtrate was concentrated, and the residue was purified by prep-HPLC (reverse phase) to give the title compound. MS (ES+): 442 (M+H) * ; (ES-): 440 (M-H) $^-$. Calc'd C₂₇H₃₁N₅O- 441.58.

Example 140

(2-{[(2-Methoxy(4-pyridy1))methyl]amino}(3-pyridyl))-N-[4-(methylethyl)phenyl]carboxamide

Step A - Preparation of (2-methoxy-4-pyridyl)methylamine

A solution of 2-methoxyisonicotinylcarboxamide (1.0 g, 6.5 mmol) and BH₃-THF complex (35 mmol) in 35 mL of THF was stirred at RT for 16 h. The reaction was quenched by addition of MeOH, and the resulting mixture was concentrated. The residue was diluted with 1N aq. NaOH and $\rm CH_2Cl_2$. The organic layer was separated, dried over $\rm Na_2SO_4$, and concentrated.

Step B - Preparation of (2-{[(2-methoxy(4-pyridyl))methyl}amino)(3-pyridyl))-N-[4-(methylethyl)phenyl]carboxamide

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The title compound was prepared from (2-methoxy-4-pyridyl)methylamine (Step A) by the method described in Example 82. MS (ES+): 377 (M+H)*; (ES-): 375 (M-H)⁻. Calc'd C₂₂H₂₄N₄O₂- 376.46.

Example 141

10 N-(3-[3-(Dimethylamino)propyl]-5-(trifluoromethyl)phenyl)-{2-[(4-pyridylmethyl)amino](3-pyridyl))carboxamide

Step A - Preparation of {3-[3-amino-5-

(trifluoromethyl)phenyl]propyn-2-yl}dimethylamine

- A mixture of 3-bromo-5-trifluoromethylaniline (1.4 g, 5.9 mmol), 1-dimethylamino-2-propyne (1.3 mL, 0.76 mmol), $PdCl_2(PPh_3)_2$ (0.26 g, 0.29 mmol) and CuI (114 mg, 0.60 mmol) in 10 mL of TEA was heated at $100^{\circ}C$ in a sealed tube for 3 h. The resulting mixture was filtered over Celite*. The filtrate was concentrated, and the residue was purified by
- prep-HPLC (reverse phase) to give the titled compound. MS (ES+): 243 (M+H)*; (ES-): 241 (M-H)⁻. Calc'd C₁₂H₁₃F₃N₂ 242.24.
- 25 <u>Step B Preparation of (3-[3-amino-5-</u> (trifluoromethyl)phenyl]propyl)dimethylamine

A mixture of the propynyl-aniline (7 g, 29 mmol, Step A) and $Pd(OH)_2$ (0.5 g) in MeOH (250 mL) was stirred under 50 psi H_2 . After 2 h, the resulting mixture was filtered over Celite. The filtrate was concentrated, and the residue was diluted with aq. 1N HCl. The aq. layer was washed with Et₂O, made basic with aq. 5N NaOH, and extracted with CH_2Cl_2 . The organic solution was dried over NaSO₄ and concentrated to give the titled compound. MS (ES+): 386 (M+H)⁺; (ES-): 384 (M-H)⁻. Calc'd $C_{18}H_{19}ClF_{3}N_{3}O$ - 385.81.

Step C - Preparation of N-(3-[3-(dimethylamino)propyl]-5(trifluoromethyl)phenyl)-{2-{(4-pyridylmethyl)amino}(3pyridyl)}carboxamide

The title compound was prepared by the method described in Example 82. MS (ES+): 458(M+H)'; (ES-): 456(M-H)'. Calc'd C₂₄H₂F₁N₅O- 457.497.

Example 142

N-[4-(tert-Buty1)-3-(3-piperidylpropy1)pheny1]{2-[(4-pyridylmethy1)amino](3-pyridy1)}carboxamide

25 Step A - Preparation of 1-piperidylprop-2-en-1-one

To a 0°C solution of acryloyl chloride (4.576 g, 50.558 mmol) in 50 ml of CH_2Cl_2 was added dropwise and very carefully piperidine (4.305 g, 50.558 mmol). The reaction flask was vented during the exothermic addition. After the

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solid.

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addition was completed, the white slurry was stirred at 0°C for 40 min and at RT for 1 h. The reaction was diluted with 70 ml CH₂Cl₂ and washed first with about 60 ml 2N HCl and then with about 60 ml of a mix of 2N NaOH and brine. The organic layer was dried over Na₂SO₄. The solution was evaporated by heating in a H₂O bath at 60°C without vacuum. Once most solvent had been evaporated off, it was furthered dried to a clear oil under high vacuum at RT for 30 min.

10 Step B - Preparation of 1-bromo-2-(tert-butyl)-5-nitrophenyl

 Br_2 (17.4 ml) was added dropwise over 40 min to a stirred mixture of 4-tert-butylnitrobenzene (59.5 g, 332 mmol), $AgSO_4$ (56.5 g, 181 mmol), H_2SO_4 (300 ml), and H_2O (33 ml) at RT. The mixture was stirred for 3 h, then poured into 0.1 M $Na_2S_2O_5/H_2O$ (1 L). The solid was filtered, washed with H_2O , Et_2O , and CH_2Cl_2 . The filtrate layers were separated. The aqueous fraction was extracted with Et_2O . The combined organic layers were combined, dried over Na_2SO_4 , and concentrated in vacuo. The yellow solid was triturated with hexanes to give a pale yellow crystalline

Step C - Preparation of (2E)-3-[2-(tert-buty1)-5nitrophenyl]-1-piperidylprop-2-en-1-one

1-(tert-Butyl)-2-bromo-4-nitrobenzene (6.885 g, 26.674 mmol, Step B), 1-piperidylprop-2-en-1-one (4.827 g, 34.677 mmol, Step A), and TEA (7.44 ml, 53.35 mmol) were dissolved into toluene (70 ml). To this solution was added Pd(OAc) $_2$ (60 mg, 0.267 mmol) and Pd(PPh $_3$) $_4$ (617 mg, 0.5335 mmol).

30 The mix was degassed with N_2 and heated in a sealed vessel at 120°C for 15 h. The reaction mixture was cooled to RT, filtered, and concentrated in vacuo. The dark crude oil was eluted through a silica gel column with 15% to 22%

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EtOAc/hexanes gradient system to yield a thick amber oil as

Step D - Preparation of (2E)-3-[2-(tert-buty1)-5-

aminophenyl]-1-piperidylprop-2-en-1-one

 $(2E)-3-[2-(tert-Buty1)-5-nitropheny1]-1-piperidy1prop-2-en-1-one (3.22 g, 10.177 mmol, step C) was dissolved in dioxane (20 ml) and IpOH (40 ml). To the <math>N_2$ -degassed solution was added 10% by weight Pd/C catalyst (2 g). The mix was placed into a Parr hydrogenator and stirred for 18 h under 60 psi H_2 . The reaction was not complete the next day, so the reaction was continued for an additional 20 h with fresh catalyst. The mix was filtered through Celite* and concentrated in vacuo to give a foamv oil.

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Step E - Preparation of 4-(tert-buty1)-3-(3piperidvlpropy1)phenylamine

(2E)-3-[2-(tert-Butyl)-5-aminophenyl]-1-piperidylprop-2-en-1-one (2.312 g, 7.619 mmol, step D) was dissolved in THF (100 ml) at RT. To this solution was added LiAlH₄ (434 mg, 11.43 mmol). After the reaction mixture stopped exotherming, it was heated at reflux at about 80°C for 4 h. The reaction was cooled to 0°C and treated by dropwise addition of 0.458 ml H₂O, 0.730 ml 10% aqueous NaOH, and 1.19 ml H₂O, respectively. The mix was stirred at RT for 40 min. Na₂SO₄ (3 g) was added and the mix was stirred for 20 min. The mix was filtered through Celite® and concentrated in vacuo. The crude was eluted through silica gel column with a gradient system of 95:5 to 90:10 CH₂Cl₂:MeOH, to yield an amber thick oil as the title compound.

Step F - Preparation of N-[4-(tert-butyl)-3-(3piperidylpropyl)phenyl]{2-[(4-pyridylmethyl)amino](3pyridyl))carboxamide

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The title compound was prepared from 4-(tert-butyl)-3-(3-piperidylpropyl)phenylamine (Step E) similar to the method described in Example 82. MS: 486.2 (M+1). Calc'd. for $C_{10}H_{19}N_{5}O$ - 485.68.

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Example 143

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N-[4-(tert-Butyl)-3-(3-pyrrolidinylpropyl)phenyl]{2-[(4-pyridylmethyl)amino](3-pyridyl)}carboxamide

MS: 472.5 (M+1). Calc'd. for $C_{29}H_{37}N_5O$ - 471.65.

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Example 144

N-[3-((1E)-4-Pyrrolidinylbut-1-enyl)-4-(tertbutyl)phenyl](2-[(4-pyridylmethyl)amino](3pyridyl))carboxamide A-733A - 276 -

MS: 484.0 (M+1). Calc'd. for C30H37N5O - 483.66.

Example 145

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N-[4-(tert-Buty1)-3-(3-morpholin-4-ylpropy1)pheny1](2-[(4-pyridylmethy1)amino](3-pyridy1))carboxamide

10 MS: 488.4 (M+1). Calc'd. for C29H37N5O2 - 487.65.

Example 146

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N-[1-(2-Morpholin-4-ylethyl)indo1-6-yl]{2-[(4-pyridylmethyl)amino](3-pyridyl)}carboxamide

Step A - Preparation of 1-(2-morpholin-4-ylethyl)indole-6ylamine

 K_2CO_3 (5.08 g, 36.726 mmol) was added to a slurry of 6-nitroindole (1.985 g, 12.242 mmol), 4-(2-chloroethyl)morpholine hydrochloride (2.278 g, 12.242 mmol),

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and CH₃CN (100 ml). The mix was heated at reflux for 18 h, then cooled to RT, filtered, and concentrated in vacuo. The crude was eluted through a silica gel column with a gradient of 3:97 to 5:95 and finally 8:92 MeOH:CH₂Cl₂, to yield upon drying 1-(2-morpholin-4-yl-ethyl)-6-nitro-1H-indole which was hydrogenated at regular condition described early to yield the title compound.

Step B - Preparation of N-[1-(2-morpholin-4-ylethyl)indol-6vl](2-[(4-pyridylmethyl)amino](3-pyridyl))carboxamide

The title compound was prepared from 1-(2-morpholin-4-ylethyl)indole-6-ylamine (Step A) similar to the method described in Example 82. MS: 457.3 (M+1). Calc'd. for $C_{26}H_{28}N_6O_2$ - 456.55.

Example 147

20 N-[4-(tert-Butyl)phenyl] {2-[(pyrimidin-4-ylmethyl)amino](3-pyridyl)}carboxamide

Step A - Preparation of pyrimidine-4-yl formaldehyde Pyrimidine-4-yl formaldehyde was prepared from 4-

25 methyl pyrimidine through a reference described in M.C. Liu et al., J Med Chem., 1995, 38 (21), 4234-4243.

Step B - Preparation of N-[4-(tert-butyl)phenyl][2[(pyrimidin-4-ylmethyl)amino](3-pyridyl)]carboxamide

The title compound was prepared from pyrimidine-4-yl formaldehyde (Step A) similar to the method described in $5 \quad \text{Example 82. MS (ES+): } 362\,\text{(M+H); (ES-): } 360\,\text{(M-H). Calc'd. for } \\ C_{21}H_{21}N_{5}O-361.19.$

Example 148

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N-(4-Chlorophenyl) {2-[(pyrimidin-4-ylmethyl)amino](3pyridyl))carboxamide

15~ MS (ES+): 340 (M+H); (ES-): 338 (M-H). Calc'd. for

 $C_{17}H_{14}C1N_5O - 339.09$.

Example 149

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{2-[(Pyrimidin-4-ylmethyl)amino](3-pyridyl)}-N-[3 (trifluorom thyl)phenyl]carboxamid

MS (ES+): 374 (M+H); (ES-): 372 (M-H). Calc'd. for $C_{18}H_{14}F_3N_5O$ 5 - 373.12.

Example 150

N-[4-(Isopropyl)phenyl] {4-[(4-pyridylmethyl)amino]pyrimidin-5-yl}carboxamide

Step A - Preparation of ethyl 2-methylthio-4-

15 [benzylamino]pyrimidine-5-carboxylate

A solution of ethyl 4-chloro-2-methylthio-pyrimidine-5-carboxylate (2.8 g, 12.2 mmol) and 4-aminomethylpyridine (1.24 mL, 12.2 mmol) in EtOH (20 mL) was heated at 70°C for 2 h. The resulting suspension was concentrated, and the residue was purified by SiO₂ chromatography to give ethyl 2-methylthio-4-[benzylamino]pyrimidine-5-carboxylate. MS (ES+): 305 (M+H)*; (ES-): 303 (M-H)⁻. Calc'd C₁₅H₁₇N₅O₂S: 303.38.

25 Step B - Preparation of N-[4-(isopropyl)phenyl](2-methylthio-4-[(4-pyridylmethyl)amino)pyrimidin-5-yl)carboxamide

- 280 -

To a solution of ethyl 2-methylthio-4-[benzylamino]pyrimidine-5-carboxylate (0.1 g, 0.3 mmol, Step A) in EtOH
(3 mL) was added 1 mL of aq. 1N NaOH solution. The resulting
mixture was stirred at 45°C for 2 h. The resulting mixture

5 was neutralized with aq. 1N HCl and concentrated. To the
residue in 3 mL of CH₂Cl₂ was added 4-isopropylaniline (90
mg, 0.66 mmol), HATU (0.18 g, 0.45 mmol), and 0.5 mL of TEA
(0.36 g, 3.5 mmol). The resulting mixture was stirred at RT
for 4 h and diluted with CH₂Cl₂. The organic solution was

10 washed with H₂O, dried over Na₂SO₄ and concentrated. The
residue was purified by SiO₂ chromatography to give N-[4(isopropyl)phenyl](2-methylthio-4-[(4pyridylmethyl)amino]pyrimidin-5-yl)carboxamide. MS (ES+):
394 (M+H)*; (ES-): 392 (M-H)*. Calc'd C₂₁H₂₂N₂OS - 393.51.

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Step C - Preparation of N-[4-(isopropyl)phenyl]{4-[(4-pyridylmethyl)amino]pyrimidin-5-yl)carboxamide

A mixture of N-[4-(isopropyl)phenyl](2-methylthio-4[(4-pyridylmethyl)amino]pyrimidin-5-yl)carboxamide (50 mg,
20 0.13 mmol, Step B) and Raney-Ni in EtOH (10 mL) was heated
at reflux for 2 h. The resulting mixture was filtered, and
the filtrate was concentrated to give the titled compound.
MS (ES+): 348 (M+H)*; (ES-): 346 (M-H)*. Calc'd C₂₀H₂₁N₅O347.42.

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Exampl 151

(2-{[(2-{2-[2-(Dimethylamino)ethoxy]ethoxy}(4pyridyl))methyl]amino}(3-pyridyl))-N-[4-(tertbutvl)phenyl]carboxamide

10 Step A - Preparation of 2-{2-[2-

$\underline{(\texttt{dimethylamino})\,\texttt{ethoxy}}\\ \underline{\texttt{pyridine-4-carbonitrile}}$

To a DMF (30 mL) solution of 2-[2-

(dimethylamino)ethoxy]ethan-1-ol (3.33 g, 25 mmol) was added NaH (60% in mineral oil, 900 mg, 22.5 mmol, hexane washed)

- and heated at 50°C for 2 h. The warm sodium alkoxide solution was added to 2-chloro-4-cyanopyridine (3.12 g, 22.5 mmol) in DMF (10 mL). After the addition, the reaction mixture was heated to 70° C for 2 h, then DMF was removed in vacuo. The residue was partitioned between $\text{CH}_2\text{Cl}_2/\text{H}_2\text{O}$. The
- O organic layer was washed with brine, dried over Na₂SO₄, and concentrated in vacuo to give a light yellow oil (5.6 g).
 MS: 236 (M+1). Calc'd. for C₁₂H₁₇N₃O₂ 235.29.

Step B - Preparation of (2-{2-[4-(aminomethyl)(2-

25 pyridyloxy)]ethoxy)ethoxy)dimethylamine

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2-{2-[2-(Dimethylamino)ethoxy]ethoxy}pyridine-4carbonitrile (330 mg 1.4 mmol, Step A) was dissolved in EtOH (10 mL) along with TEA (2 mL) and suspended with Pd/C (10%, 40 mg). The reaction mixture was stirred overnight at RT under balloon filled with Ho. After removing the balloon. the reaction suspension was filtered through a layer of Celite®. The Celite® layer was rinsed with MeOH. The combined filtrate was concentrated in vacuo to give a light vellow oil. MS: 240 (M+1). Calc'd. for C12H21N3O2 - 239.32.

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Step C - Synthesis of 2-fluoropyridine-3-carboxylic acid

To a solution of 2-fluoropyridine (10 g, 100 mmol) in THF (150 mL) under -78°C was dropwise added an LDA solution (2M in heptane/THF/ethylbenzene, 60 mL). The mixture was stirred at -78°C for 3 h after the addition of LDA then 15 quenched with No dried solid COo. After warming to RT, the reaction was partitioned between EtOAc (100 mL) and H2O (200 mL). The aqueous layer was acidified to pH between 3-4 and extracted with EtOAc. The organic solution was collected and washed with brine and dried over Na2SO4. After removing solvent in vacuum, a brown oil was received as the desired compound. MS: 140 (M-H). Calc'd. for C6H4FNO2 - 141.10.

Step D - Synthesis of 2-fluoropyridine-3-carbonyl chloride

2-Fluoropyridine-3-carboxylic acid (7 g, Step C) was suspended in SOCl2 (100 mL). After heating under reflux for 2 h, the mixture became homogeneous. Excess SOCl2 was removed in vacuo to afford a brown solid as desired product.

3.0 Step E - Synthesis of N-[4-(tert-butyl)phenyl] 2fluoropyridine-3-carboxamide

To a suspension of 2-fluoropyridine-3-carbonyl chloride (3.2 g. 20 mmol, Step D) and NaHCO3 (4 g. 48 mmol) in CH2Cl2 added in dropwise a solution of 4-tert

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Calc'd, for C28H37N5O3 - 491.63

butylaniline (3.0 g, 20 mmol). After the addition, the suspension was stirred at RT for 5 h. Solid inorganic salts were removed via filtration. The filtrate was concentrated to afford a brown solid as desired compound. MS: 273 (M+H). Calc'd. for $C_{14}H_1 \ge FN_2O - 272.33$.

Step F - Synthesis of (2-[({2-[2-(2-N,N-dimethylaminoethoxy]ethoxy]-4-pyridyl)methyl)amino](3-pyridyl))-N-(4-tert-butylphenyl)carboxamide
N-[4-(tert-Butyl)phenyl] 2-fluoropyridine-3-

carboxamide (544 mg, 2 mmol, Step E) was dissolved in pyridine (5 mL) along with (2-(2-[4-(aminomethyl) (2-pyridyloxy)]ethoxy)ethoxy)dimethylamine (570 mg, 2.38 mmol, Step A). The reaction was heated to 85°C for 48 h. After removal of pyridine in vacuo, the residue was dissolved in CH_2Cl_2 and washed with NaHCO₃ (Sat. aq), then brine. After drying over Na₂SO₄, the CH_2Cl_2 solution was concentrated in vacuo and purified via prep. HPLC (H_2O/CH_3CN : 5%-95% gradient) to give the title product. MS: 492 (M+1).

The following compounds (Examples 152-157) were analogously synthesized by the method described in Example 151 unless specifically described. Detailed intermediate preparations are included.

Example 152

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{2-[(4-Pyridylmethyl)amino](3-pyridyl)}-N-{4-[2,2,2trifluoro-1-(2-piperidylethoxy)-1(trifluoromethyl)ethyl]phenyl)carboxamide

10 Step A - Preparation of 4-[2,2,2-trifluoro-1-(2-piperidin-1-yl-ethoxy)-1-trifluoromethyl-ethyl]-phenylamine

DEAD (366 mg, 2.1 mmol) was added drop-wise to the

solution of 2-(4-aminophenyl)-1,1,1,3,3,3-hexafluoropropan-2-ol (520 mg, 2 mmol), 2-piperidylethan-1-ol (260 mg, 2 mmol) and PPh₃ (550 mg, 2.1 mmol) in THF (10 mL). The mixture was stirred for 2 h. The reaction was partitioned between EtOAc and aqueous NaHCO, solution and the organic phase was washed with brine. After concentrated in vacuo, the organic residue was purified by flash chromatography on silica to give the title intermediate.

Step B - Preparation of {2-[(4-pyridylmethyl)amino](3pyridyl)}-N-(4-[2,2,2-trifluoro-1-(2-piperidylethoxy)-1-(trifluoromethyl)ethyl]phenyl)carboxamide

The title compound was synthesized by the method described in Example 151. MS: 582 (M+1). Calc'd. for $C_{28}H_{22}F_4N_5O_2 \ - \ 581.56 \ .$

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Example 153

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(2-{[(2-{2-[2-(Dimethylamino)ethoxy]ethoxy}(4pyridyl))methyl]amino)-6-fluoro(3-pyridyl))-N-[3(trifluoromethyl)phenyl]carboxamide

10 Step A Preparation of 2,6-difluoropyridine-3-carbonyl chloride

The title compound was prepared similar to that described in Example 151, Step D.

15 Step B - Preparation of (2-[[(2-(2-[2-(dimethylamino)-ethoxy]ethoxy)(4-pyridyl))methyl]amino)-6-fluoro(3-pyridyl))-N-[3-(trifluoromethyl)phenyl]carboxamide

The title compound was synthesized by the method described in Example 151. MS: 522 (M+1). Calc'd. for $C_{25}H_{27}F_4N_5O_3$ - 521.51.

ADDESCE DESCRIPTION

Example 154

N-[4-(tert-Butyl)phenyl]{6-fluoro-2-[(4pyridylmethyl)amino](3-pyridyl)}carboxamide

Step A - Preparation of N-(4-tert-butyl-phenyl)-2,6difluoro-nicotinamide

A solution of 2,6-difluoropyridine-3-carboxylic acid (3.2 g, 20 mmol), t-butylaniline (3.0 g, 20 mmol), HOBt (2.6 g, 20 mmol), EDAC (8 g, 40 mmol), and DIEA (8 mL) in $\mathrm{CH}_2\mathrm{Cl}_2$ (80 mL) was stirred at RT for 1 h. The mixture was washed with aq. NaHCO3 and brine. The organic solution was dried over Na₂SO₄ and concentrated in vacuo. The residue was purified via flash chromatography on silica (Hex:EtOAc = 4:1) to give a light yellow flaky crystal as desired product.

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Step B - Preparation of N-[4-(tert-butyl)phenyl] (6-fluoro-2-[(4-pyridylmethyl)amino](3-pyridyl))carboxamide

The title compound was synthesized similar to that described in Example 151 except that it was synthesized at RT. MS: 379 (M+1). Calc'd. for $C_{22}H_{23}FN_4O$ - 378.45.

The following compounds were analogously synthesized by the method described in Example 154. Detailed intermediate preparations are described.

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Example 155

{6-Fluoro-2-[(4-pyridylmethyl)amino](3-pyridyl)}-N-[4-(isopropyl)phenyl]carboxamide

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MS: 365 (M+1). Calc'd. for C21H21FN4O - 364.42.

Example 156

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{6-Fluoro-2-[(4-pyridylmethyl)amino](3-pyridyl)}-N-[3-(trifluoromethyl)phenyl]carboxamide

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MS: 391 (M+1). Calc'd. for $C_{19}H_{14}F_4N_4O$ - 390.34.

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Example 157

5 N-(1-Bromo(3-isoquinoly1))(6-fluoro-2-[(4-pyridy1methy1)amino](3-pyridy1)}-carboxamide

 $\label{eq:ms:def} \text{MS:} \quad 452/454 \text{ (M+1).} \quad \text{Calc'd. for} \quad C_{21}H_{15}\text{BrFN}_5\text{O} \text{ - } 452.29\,.$

Example 158

N-(4-Phenoxyphenyl) {2-[(4-pyridylmethyl)amino](3pyridyl)}carboxamide

Step A - Preparation of (2-chloro(3-pyridy1))-N-(4-phenoxy-pheny1)carboxamide

2-Chloronicotinic acid (0.78 g, 5.0 mmol) and TEA (1.6 20 ml, 10.0 mmol) were added to anhydrous THF (50 ml) under a N₂ atmosphere at 0°C. After stirring for 5 min, ethyl

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chloroformate (0.54 g, 5.0 mmol) was added dropwise and the mixture gradually came to RT over a period of 1 h. 4-Phenoxyaniline (0.83 g, 5.0 mmol) was added and the mixture was stirred for 14 h. The mixture was partitioned between $\rm H_{2}O$ and EtOAc. The aqueous layer was extracted two additional times with EtOAc (50 ml). The combined organic layers were then washed with brine, dried over $\rm Na_{2}SO_{4}$, and evaporated. The resulting brown oil was used directly in the subsequent reaction without further purification. MS $\rm m/z$: 325 (M + 1). Calc'd for $\rm C_{18}H_{12}ClN_{2}O_{2}$: 324.07.

Step B - Preparation of N-(4-phenoxyphenyl)(2-[(4pyridylmethyl)amino](3-pyridyl)}carboxamide hydrochloride

The amide (0.500 g, 1.5 mmol, Step A) and 415 aminomethylpyridine (0.486 g, 4.5mmol) were combined and heated neat at 90°C for 48 h. After cooling to RT, the mixture was poured into a saturated NaHCO₃ solution and extracted with EtOAc. The combined organic layers were washed with brine, dried over Na₂SO₄, and evaporated. The

- 20 resulting brown oil was purified by column chromatography with EtOAc/hexanes (2:1) as eluant to leave N-(4phenoxyphenyl){2-[(4-pyridylmethyl)amino](3pyridyl)}formamide as a clear oil. This material was converted directly into the HCl salt by dissolution in MeOH
- 25 (5 ml), treatment with 3 equivalents of an HCl ethereal solution, and evaporation of solvent to leave the titled product as a light yellow solid. MS (ES+): 397 (M+H)*; (ES-): 395 (M-H). Calc'd. for C₂₄H₂₀N₄O₂ 396.16.
- 30 The following compounds (Examples 159-161) were prepared similar to the method described in Example 158.

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Example 159

- 290 -

N-(4-Biphenyl)(2-[(4-pyridylmethyl)amino](3pyridyl))carboxamide

MS: 381 (M+1); 379 (M-1). Calc'd. for $C_{24}H_{20}N_4O$ - 380.16.

Example 160

N-(3-Phenoxyphenyl) {2-[(4-pyridylmethyl)amino](3-pyridyl)}carboxamide hydrochloride

MS: 397 (M+1); 395 (M-1). Calc'd. for $C_{24}H_{20}N_4O_2$ - 396.16.

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Example 161

5 N-(4-Cyclohexylphenyl){2-[(4-pyridylmethyl)amino](3-pyridyl)}carboxamide

MS: 387 (M+1); 385 (M-1). Calc'd. for $C_{24}H_{26}N_4O$ - 386.21.

Example 162

N-(4-Imidazol-1-ylphenyl) {2-[(4-pyridylmethyl)amino](3pyridyl))carboxamide

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Step A - Preparation of (2-Chloro(3-pyridyl))-N-(4-imidazolylphenyl)carboxamide

A slurry of 4-imidazolylphenylamine (15.9 mg, 0.100 mmol), polymer-supported DIPEA (0.100 g, 0.362 mmol, 3.62 mmol/g loading) in CH_2Cl_2 (2 ml) was treated with a 2-chloropyridine-3-carbonyl chloride solution (0.10 M, 0.200 mmol, 2.0 ml, 2.0 eg) in CH_2Cl_2 . The mixture was vortexed at RT for 14 h. Afterwards, the excess acid chloride was removed by treating the reaction mixture with polymer-

10 supported trisamine resin (0.100 g, 0.375 mmol, 3.75 mmol/g loading). The slurry was shaken at RT for an additional 18 h. The reaction mixture was filtered, rinsed with CH₂Cl₂ (1 ml), and the filtrate was concentrated under reduced pressure. The resulting brown oil was used directly in the subsequent reaction.

Step B - Synthesis of N-(4-imidazolylphenyl){2-[(4pyridylmethyl)amino](3-pyridyl)}carboxamide hydrochloride (2-Chloro-(3-pyridyl))-N-(4-imidazolylphenyl)-

20 carboxamide was treated with 4-aminomethylpyridine (0.100 g, 0.93 mmol) and heated neat at 120°C for 18 h. After cooling to RT, the material was purified by preparative HPLC. The final product was converted into an HCl salt by dissolution in a minimum of MeOH, treatment with an HCl ethereal

solution, and evaporation of solvent. MS: (ES+) 371 (M + 1)⁺; (ES-): 369 (M - 1)⁻. Calc'd. for C₂₁H₁₈N₆O - 370.15.

The following compounds (Examples 163-166) were analogously synthesized by the method described in Example 162.

Example 163

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N-(4-Morpholin-4-ylphenyl)(2-[(4-pyridylmethyl)amino](3-pyridyl))carboxamide

The title compound was isolated as the HCl salt. 10 MS: 390 (M+1); 388 (M-1). Calc'd. for $C_{22}H_{23}N_5O_2$ - 389.19.

Example 164

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N-(4-Cyanonaphthyl) {2-[(4-pyridylmethyl)amino](3pyridyl))carboxamide

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The title compound was isolated as the HCl salt. MS: 380 (M+1); 378 (M-1). Calc'd. for $C_{21}H_{17}N_5O$ - 379.14.

Example 165

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{2-[(4-Pyridylmethyl)amino](3-pyridyl)}-N-[4-(trifluoromethyl)phenyl]carboxamide

10 The title compound was isolated as the HCl salt. MS: 373 (M+1); 371 (M-1). Calc'd. for $C_{10}H_{15}F_{1}N_{4}O$ - 372.12.

Example 166

15

Methyl-({2-[(4-pyridylmethyl)amino]-3pyridyl}carbonylamino)benzoate

The title compound was isolated as the HCl salt. 20 MS: 363 (M+1); 361 (M-1). Calc'd. for $C_{20}H_{18}N_4O_3$ - 362.14. LOCHECOL CILOCO

The following compounds were synthesized by a procedure similar to the method described in Example 3, using an aldehyde to react with the aminopyridine core via reductive amination.

Example 167

10

N-[4-(Isopropy1)pheny1]{2-[(4-quinolylmethy1)amino](3pyridy1)}carboxamide

MS: (ES+) 397 (M+H); (ES-) 395 (M-H). Calc'd. for $C_{25}H_{24}N_4O$ - 15 396.20.

Example 168

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N-[4-(tert-Butyl)phenyl]{2-[(6-quinolylmethyl)amino](3-pyridyl)}carboxamide

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MS (ES+): 411 (M+H); (ES-): 409 (M-H). Calc'd. for $C_{26}H_{26}N_4O$ - 410.51.

Example 169

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{2-[(6-Quinolylmethyl)amino](3-pyridyl)}-N-[3-(trifluoromethyl)phenyl]carboxamide

10

MS (ES+): 423 (M+H); (ES-): 421 (M-H). Calc'd. for C23H17F3N4O: 422.14.

Other compounds included in this invention are set 15 forth in Tables 3-9 below.

 R^1

 \mathbb{R}^2

_	<u> </u>		
	170. 2-chlorophenyl	Н	1
	171.4-benzimidazolyl	H	1
	172.5-benzimidazolyl	Н	1
	173.7-benzimidazolyl	Н	1
10	174.2-chlorophenyl	5-Br	1
	175.3-isoquinolinyl	5-Br	1
	176.2-quinolinyl	5-Br	1
	177.2-benzthiazolyl	5-Br	1
	178.2-benzimidazolyl	5-Br	1
15	179.4-benzimidazolyl	5-Br	1
	180.5-benzimidazolyl	5-Br	1
	181.6-benzimidazolyl	5-Br	1
	182.7-benzimidazolyl	5-Br	1
	183.4-chlorophenyl	H	3
20	184.4-chlorophenyl	3-pyridyl	1
	185.4-pyridyl	Н	1
	186.4-pyridyl	6-CH ₃	1
	187.4-chlorophenyl-	5-C1	1
	188.3,4-dichlorophenyl-	5-Br	1
25	189. 4-fluorophenyl	6-CH ₃	1
	190. 3-chlorophenyl	6-CH ₃	1
	191. 3-fluorophenyl	6-CH ₃	1
	192. 3-fluoro-4-methoxyphenyl	6-CH ₃	1
	193. 3-fluoro-4-methylphenyl	6-C1	1
	15	171. 4-benzimidazolyl 172. 5-benzimidazolyl 173. 7-benzimidazolyl 10 174. 2-chlorophenyl 175. 3-isoquinolinyl 176. 2-quinolinyl 177. 2-benzthiazolyl 178. 2-benzimidazolyl 180. 5-benzimidazolyl 180. 5-benzimidazolyl 181. 6-benzimidazolyl 182. 7-benzimidazolyl 183. 4-chlorophenyl 20 184. 4-chlorophenyl 185. 4-pyridyl 186. 4-pyridyl 187. 4-chlorophenyl 188. 3,4-dichlorophenyl- 25 189. 4-fluorophenyl 190. 3-chlorophenyl 191. 3-fluorophenyl 192. 3-fluoro-4-methoxyphenyl	171. 4-benzimidazolyl H 172. 5-benzimidazolyl H 173. 7-benzimidazolyl H 173. 7-benzimidazolyl H 174. 2-chlorophenyl 5-Br 175. 3-isoquinolinyl 5-Br 176. 2-quinolinyl 5-Br 177. 2-benzthiazolyl 5-Br 178. 2-benzimidazolyl 5-Br 180. 5-benzimidazolyl 5-Br 180. 5-benzimidazolyl 5-Br 181. 6-benzimidazolyl 5-Br 182. 7-benzimidazolyl 5-Br 182. 7-benzimidazolyl 5-Br 182. 4-chlorophenyl H 20 184. 4-chlorophenyl 3-pyridyl 185. 4-pyridyl H 186. 4-pyridyl 6-CH3 187. 4-chlorophenyl- 5-Br 25 189. 4-fluorophenyl- 5-Br 26 189. 4-fluorophenyl 6-CH3 190. 3-chlorophenyl 6-CH3 191. 3-fluorophenyl 6-CH3 192. 3-fluoro-4-methoxyphenyl 6-CH3

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Table 3. (cont.)

5	#	R ¹	 R ²	n	
	194.	4-phenoxyphenyl	H	1	
	195.	3-phenoxyphenyl	H	1	
	196.	4-biphenyl	H	1	
	197.	4-cyclohexylphenyl	H	1	
10	198.	2-quinolyl	H	1	
	199.	3-isoquinolyl	H	1	
	200.	3-quinolyl	H	1	
	201.	1-isoquinolyl	H	1	
	202.	5-quinolyl	H	1	
15	203.	5-isoquinolyl	H	1	
	204.	6-quinolyl	H	1	
	205.	6-isoquinolyl	H	1	
	206.	7-quinolyl	H	1	
	207.	7-isoquinolyl	H	1	
20	208.	4-quinolyl	H	1	
	209.	4-isoquinolyl	H	1	
	210.	4-pyridyl	H	1	
	211.	4-pyrimidinyl	Н	1	
	212.	2-pyrimidinyl	H	1	
25	213.	6-pyrimidinyl	H	1	
	214.	4-pyridazinyl	H	1	
	215.	5-pyridazinyl	H	1	
	216.	4-indolyl	Н	1	
	217.	5-isoindolyl	н	1	

HOCKET OF HOOF

Table 3. (cont.)

5	#	R ¹	R ²	n
	218.	5-naphthyridinyl	Н	1
	219.	6-quinozalinyl	H	1
	220.	6-isoquinolyl	H	1
	221.	4-naphthyridinyl	H	1
10	222.	5-quinozalinyl	H	1
	223.	4-naphthyridinyl	H	1
	224.	7-tetrahydroquinolinyl	H	1
	225.	6-indazolyl	H	1
	226.	6-isoindolyl	H	1
15	227.	5-indazolyl	H	1
	228.	5-isoindolyl	H	1
	229.	6-benzothienyl	H	1
	230.	6-benzofuryl	H	1
	231.	5-benzothienyl	H	1
20	232.	5-benzofuryl	H	1
	233.	2-benzimidazolyl	H	1
	234.	2-benzoxazolyl	Н	1
	235.	2-benzthiazolyl	H	1
	236.	6-benzimidazolyl	Н	1
25	237.	6-benzoxazolyl	Н	1
	238.	6-benzthiazolyl	H	1
	239.	2-quinazolinyl	н	1
	240.	3-(phenoxy)-6-pyridyl	Н	1
	241.	4-(phenylcarbonyl)phenyl	Н	1

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Table 3. (cont.)

242. 4-(phenylamino)phenyl H	1
243. 4-cyclohexyloxyphenyl H	1
244. 4-(3-thienyl)phenyl H	1
245. 4-(pyrazol-3-yl)phenyl H	1
10 246. 4-chlorophenyl 6-F	2
247. 4-pyridyl 6-Cl	1
248. 3-methoxyphenyl 6-F	1
249. 4-hydroxyphenyl 6-Cl	1
250. 3-hydroxyphenyl H	1
15 251. 2-hydroxyphenyl H	1
252. 4-chlorophenyl 6-F	1
253. 4-phenoxyphenyl 6-F	1
254. 4-biphenyl 6-phenyl	1
255. 4-hydroxyphenyl 6-phenyl	1
20 256. 4-cyclohexylphenyl 6-F 1	
257. 3-isoquinolyl 6-phenyl	1
258. 4-piperidinylmethylphenyl H	1
259. 4-morpholinylmethylphenyl H	1

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Table 4a.

5 # R¹

260.4-chlorophenyl

261.3,4-dichlorophenyl

262.4-phenoxyphenyl

10 263.4-biphenyl

264.4-cyclohexylphenyl

265.3-isoquinolyl

15

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Table 4b.

R1

266.4-chlorophenyl

267.3,4-dichlorophenyl

268.4-phenoxyphenyl

269.4-biphenyl

25 270.4-cyclohexylphenyl

271.3-isoquinolyl

A-733A

Table 4c.

5	#	R ¹	A ⁵	
	272	. 4-chlorophenyl	NH	
		. 3,4-dichlorophenyl	NH	
	274	. 4-phenoxyphenyl	NH	
10	275	.4-biphenyl	NH	
	276	. 4-cyclohexylphenyl	NH	
	277	.3-isoquinolyl	NH	
	278	.4-chlorophenyl	О	
	279	.3,4-dichlorophenyl	0	
15	280	. 4-phenoxyphenyl	О	
	281	.4-biphenyl	0	
	282	.4-cyclohexylphenyl	0	
	283	.3-isoquinolyl	0	
	284	.3,4-dichlorophenyl	s	
20	285	. 4-phenoxyphenyl	S	
	286	.4-biphenyl	S	
	287	. 4-cyclohexylphenyl	S	
	288	.3-isoguinolyl	S	

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Table 4d.

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# R ¹	A^5
289.4-chlorophenyl	NH
290.3,4-dichlorophenyl	NH
291.4-phenoxyphenyl	NH
292.4-biphenyl	NH
293.4-cyclohexylphenyl	NH
294.3-isoquinolyl	NH
295.4-chlorophenyl	0
296.3,4-dichlorophenyl	0
297.4-phenoxyphenyl	0
298.4-biphenyl	o
299. 4-cyclohexylphenyl	0
300.3-isoquinolyl	0
301.3,4-dichlorophenyl	s
302.4-phenoxyphenyl	s
303.4-biphenyl	s
304.4-cyclohexylphenyl	S
305.3-isoquinolyl	s

Table 4e.

#	R ¹	A ⁵	
306	. 4-chlorophenyl	NH	
307	.3,4-dichlorophenyl	NH	
308	3.4-phenoxyphenyl	NH	
309	. 4-biphenyl	NH	
310	. 4-cyclohexylphenyl	NH	
311	3-isoquinolyl	NH	
312	. 4-chlorophenyl	0	
313	.3,4-dichlorophenyl	0	
314	. 4-phenoxyphenyl	0	
315	. 4-biphenyl	0	
316	. 4-cyclohexylphenyl	0	
317	. 3-isoquinolyl	0	
318	3.3,4-dichlorophenyl	s	
319	. 4-phenoxyphenyl	s	
320	. 4-biphenyl	s	
321	4-cyclohexylphenyl	s	
322	.3-isoquinolyl	s	

10

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337.4-biphenyl

339.3-isoquinolyl

338. 4-cyclohexylphenyl

Table 4f.

s

s

A⁵ 323. 4-chlorophenyl NH 324.3,4-dichlorophenyl NH 325. 4-phenoxyphenyl NH 326.4-biphenyl NH 327.4-cyclohexylphenyl NH NH 328.3-isoquinolyl 329. 4-chlorophenyl 0 330.3,4-dichlorophenyl 0 331.4-phenoxyphenyl 0 332.4-biphenyl 0 333.4-cyclohexylphenyl 0 334.3-isoquinolyl 0 335.3,4-dichlorophenyl S 336.4-phenoxyphenyl s

Table 4g.

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# R ¹	A ⁵
340.4-chlorophenyl	NH
341.3,4-dichlorophenyl	NH
342.4-phenoxyphenyl	NH
343.4-biphenyl	NH
344.4-cyclohexylphenyl	NH
345.3-isoquinolyl	NH
346.4-chlorophenyl	0
347.3,4-dichlorophenyl	0
348.4-phenoxyphenyl	0
349.4-biphenyl	0
350.4-cyclohexylphenyl	0
351.3-isoquinolyl	0

20

15

Table 4h.

	,	# R	A ⁵
		352.4-chlorophenyl	NCH ₃
		353.3,4-dichlorophenyl	NCH ₃
		354.4-phenoxyphenyl	NCH ₃
j (2		355.4-biphenyl	NH
9	10	356.4-cyclohexylphenyl	NH
F O		357.4-tert-butylphenyl	NCH ₃
(f). (0		358.4-chlorophenyl	0
ķd		359.3,4-dichlorophenyl	0
: (2		360.4-phenoxyphenyl	0
hab :	15	361.4-biphenyl	0
		362.4-cyclohexylphenyl	0
and a		363.3-isoquinolvl	0

Table 5.

$$R^2 \xrightarrow{\begin{array}{c} 5 \\ 5 \\ 4 \end{array}} \stackrel{\bigcirc}{\underset{R}{|}} N^{-R^1}$$

5	#	R	Y	R ¹	R ²
	364.	4-pyridyl	-NHSO ₂ -	4-chlorophenyl	н
	365.	4-pyridyl	-NHSO2-	4-chlorophenyl	5-Br
	366.	4-pyridyl	-NHSO ₂ -	3-chlorophenyl	Н
10	367.	4-pyridyl	-NHSO ₂ -	3-chlorophenyl	5-Br
	368.	4-pyridyl	-NHSO ₂ -	4-phenoxyphenyl	Н
	369.	4-pyridyl	-NHSO ₂ -	4-biphenyl	Н
	370.	4-pyridyl	-NHSO ₂ -	3-isoquinolyl	H
	371.	4-pyridyl	-NHSO ₂ -	3-isoquinolyl	5-Br
15	372.	5-quinolyl	-NHSO ₂ -	4-chlorophenyl	Н
	373.	5-quinolyl	-NHSO ₂ -	4-chlorophenyl	5-Br
	374.	5-quinolyl	-NHSO ₂ -	3-chlorophenyl	Н
	375.	5-quinolyl	-NHSO ₂ -	3-chlorophenyl	5-Br
	376.	5-quinolyl	-NHSO ₂ -	4-phenoxyphenyl	Н
20	377.	6-quinolyl	-NHSO ₂ -	4-biphenyl	H
	378.	5-quinolyl	-NHSO ₂ -	3-isoquinolyl	Н
	379.	6-quinolyl	-NHSO ₂ -	3-isoquinolyl	5-Br
				F F F	
	380.	4-pyridyl	-NHCH ₂ -	F	Н
	381.	CH ₃	-NHCH ₂ -	HN-N	н
25					

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Table 5. cont.

5	#	R	Y	R ¹	R ²
	382.	H O CH ₃	-NHCH ₂ -	HN-N	Н
	383.	N CH3	-NHCH ₂ -	HN-N F	Н
	384.	N CH ₃	-NHCH ₂ -	$\bigcap_{F} \bigcap_{F} F$	Н
	385.	N CH3	-NHCH ₂ -	4-CF,-phenyl	Н
10	386.	N O CH3	-NHCH ₂ -	- N	н
	387.	N CH3	-NHCH ₂ -	HN-N F	Н
	388.	N O CH3	-NHCH ₂ -	$\bigvee_{F} \bigvee_{F}^{F}$	Н
	389.	N O CH ₃	-NHCH ₂ -	3-CF ₃ -phenyl	Н
	390.	N O N CH3	-NHCH ₂ -	$-\bigcirc\!$	Н

Table 5. cont.

#	R	Y	R ¹	R ²
391.	N CH3	-NHCH ₂ -	H ₃ C CH ₃	Н
392.	N CH3	-NHCH ₂ -	4-CF,-phenyl	Н
393.	ŗ ^N Ţ ⁰ ∨ CH³	-NHCH ₂ -	-N	Н
394.	N CH ³	-NHCH ₂ -	HN-N F	н
395.	N CH3	-NHCH ₂ -	F_{F}	н
396.	µ ^{CH³}	-NHCH ₂ -	3-CF ₃ -phenyl	Н
397.	N CH3	-NHCH ₂ -	H ₀ C	Н
398.	N CH3	-NHCH ₂ -	H ₃ C CH ₃	Н

Table 5. cont.

$$R^{2} \underbrace{ \left(\begin{array}{c} 0 \\ 5 \\ 4 \\ 3 \\ 2 \end{array} \right)}_{R} N^{-R^{1}}$$

			K	
#	R	Y	R ¹	R ²
399.	N CH3	-NHCH ₂ -	S-N	Н
400.	N CH3	-NHCH ₂ -	4-CF ₃ -phenyl	Н
401.	N CH3	-NHCH ₂ -	HN-N F	Н
402.	N CH3	-NHCH ₂ -	$\bigcap_{F} \bigcap_{F}^{F}$	Н
403.	CN CH3	-NHCH ₂ -	3-CF ₃ -phenyl	Н
404.	N CH3	-NHCH ₂ -	H ₃ C CH ₃	Н
405.	N CH ₃	-NHCH ₂ -	N H	Н
406.	N CH3	-NHCH ₂ -	4-CF ₃ -phenyl	Н
407.	N N N N N N N N N N N N N N N N N N N	-NHCH ₂ -	-N	Н

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5

Table 5. cont.

$$\mathbb{R}^2 \underbrace{ \left(\begin{array}{c} 0 \\ 5 \\ 4 \\ 3 \end{array} \right) }_{N} \mathbb{R}^1$$

#	R	Y	R ¹	R ²
408.	N N N N N N N N N N N N N N N N N N N	-NHCH ₂ -	HN-N	н
409.	N NINHE	-NHCH ₂ -	F	н
410.	N NIMe	-NHCH ₂ -	3-CF,-phenyl	н
411.	N NMe	-NHCH ₂ -	H ₃ Ç CH ₂	н
412.	N NOME	-NHCH ₂ -	N H	н
413.	N NMe	-NHCH ₂ -	4-CF,-phenyl	н
414.	4-pyridyl	-NHCH ₂ -	N N	н
415.	4-pyridyl	-NHCH ₂ -		н
416.	4-pyridyl	-NHCH ₂ -		н

Table 5. cont.

$$\mathbb{R}^2 \underbrace{ \left(\begin{array}{c} 0 \\ 5 \\ 4 \\ 3 \end{array} \right) }_{\mathbb{R}} \mathbb{R}^1$$

			R	
#	R	Y	R ¹	R ²
417.	4-pyridyl	-NHCH ₂ -	N N N	н
418.	4-pyridyl	-NHCH ₂ -	N NH	Н
419.	4-pyrimidinyl	-NHCH ₂ -	N N N	Н
420.	4-pyrimidinyl	-NHCH ₂ -	NH NH	н
421.	4-pyrimidinyl	-NHCH ₂ -		Н
422.	4-pyrimidinyl	-NHCH ₂ -	N N	н
423.	4-pyrimidinyl	-NHCH ₂ -		Н

Table 5. cont.

			-	•	
5	#	R	Y	R ¹	R ²
	424.	4-pyrimidinyl	-NHCH ₂ -		н
	425.	4-pyrimidinyl	-NHCH ₂ -	N CH,	Н
	426.	4-pyrimidinyl	-NHCH ₂ -		Н
	427.	4-pyrimidinyl	-NHCH ₂ -	N~CH ₃	н
10	428.	4- pyrimidinyl	NHCH ₂ -		Н
	429.	3-pyridyl	-NH (CH ₂) ₂ -		н
	430.	3-pyridyl	-NH (CH ₂) ₂ -	M N N	н

Table 5. cont.

$$\begin{array}{c|c} \mathbb{R}^2 & \bigcirc & \mathbb{R}^1 \\ & \searrow & \mathbb{R}^1 \\ & & \mathbb{R} \end{array}$$

			I	ŧ.	
5	#	R	Y	R ¹	R ²
	431.	3-pyridyl	-NH (CH ₂) ₂ -		н
	432.	3-pyridyl	-NH (CH ₂) ₂ -		Н
	433.	3-pyridyl	-NH (CH ₂) ₂ -	N CH,	н
	434.	3-pyridyl	-NH (CH ₂) ₂ -		н
.0	435.	3-pyridyl	-NH (CH ₂) ₂ -	N-CH3	н
	436.	3-pyridyl	-NH (CH ₂) ₂ -	NH NH	Н
	437.	3-pyridyl	-NH (CH ₂) ₂ -	N N N	н

Table 5. cont.

$$\mathbb{R}^2 \underbrace{\begin{array}{c} 0 \\ 5 \\ 4 \\ 3 \\ N \end{array}}_{N} \mathbb{R}^1$$

			R	
#	R	Y	R ¹	R ²
438.	OMe	-NHCH ₂ -	N N N	Н
439.	OMe	-NHCH ₂ -	N NH	Н
440.	OMe	-NHCH ₂ -		н
441.	OMe	-NHCH ₂ -	N-c	н _{3 Н}
442.	OMe	-NHCH ₂ -		Н
443.	OMe	-NHCH ₂ -		н
444.	T NOME	-NHCH ₂ -		Н

Table 5. cont.

$$\mathbb{R}^2 \underbrace{ \left(\begin{array}{c} 0 \\ 5 \\ 3 \end{array} \right) \left(\begin{array}{c} 0 \\ N \end{array} \right)}_{I} \mathbb{R}^1$$

#	R	Υ	R ¹	R ²
445.	OMe	-NHCH ₂ -		Н
446.	OMe	-NHCH ₂ -	N _{CH3}	Н
447.	SMe	-NHCH ₂ -		н
448.	SMe	-NHCH ₂ -	N N N	н
449.	SMe	-NHCH ₂ -		Н
450.	\bigvee_{N} SMe	-NHCH ₂ -		Н
451.	\bigvee_{N}^{SMe}	-NHCH ₂ -	N CH ₃	Н
452.	CH_3	-NHCH ₂ -		н

10

Table 5. cont.

#	R	Y	R ¹	R ²
453.	CH ₃	-NHCH ₂ -	N N	н
454.	CH ₃	-NHCH ₂ -		н
455.	CH ₃	-NHCH ₂ -		н
456.	CH ₃	-NHCH ₂ -	N CH ₃	н
457.	N (CH ₃)	-NHCH ₂ -		н
458.	N (CH ₃)	-NHCH ₂ -	N N N	н
459.	N(CH ₃)	-NHCH ₂ -		н
460.	N (CH ₃)	-NHCH ₂ -		н

Table 5. cont.

$$R^2 \underbrace{ \left(\begin{array}{c} 4 \\ 5 \\ 6 \\ N \end{array} \right)^2}_{I} \underbrace{ \left(\begin{array}{c} 0 \\ N \end{array} \right)^{-R^1}}_{I}$$

10

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#	R	Y	R ¹	R ²
461.	$\bigvee_{N}^{N(CH_3)_2}$	-NHCH ₂ -	N _{CH₃}	н
462.	4-pyridyl	-NHCH ₂ -	N CH ₃	Н
463.	4-pyridyl	-NHCH ₂ -		н
464.	4-pyridyl	-NHCH ₂ -	CF ₂ CF ₃ N-CH ₃	н
465.	4-pyridyl	-NHCH ₂ -	N—H	Н
466.	4-pyridyl	-NHCH ₂ -	CF ₂ CF ₃	Н
467.	4-pyridyl	-NHCH ₂ -	N—CH ₃	Н
468.	4-pyridyl	-NHCH ₂ -	H N	Н

Table 5. cont.

$$R^2 \xrightarrow{\begin{cases} 5 & 4 & 3 \\ 6 & 2 \end{cases}} N^{R^1}$$

 \mathbb{R}^1 \mathbb{R}^2 Y 469. 4-pyridyl -NHCH₂-Н CF2CF3 470. 3-pyridyl -NH (CH₂)₂-Н 471. 3-pyridyl -NH (CH₂)₂-Н -NH (CH₂)₂-472. 3-pyridyl н 10 473. 3-pyridyl -NH (CH₂)₂н 474. 3-pyridyl -NH(CH₂)₂-Н -NH (CH₂)₂-475. 3-pyridyl Н 476. 3-pyridyl -NH (CH₂)₂-Н

Н

15 477. 3-pyridyl -NH(CH₂)₂-

10

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Table 5. cont.

 R^2 R 478. 3-pyridyl -NH (CH₂)₂-Н 479. 4-pyrimidinyl -NHCH2-Н 480. 4-pyrimidinyl -NHCH2-Н 481. 4-pyrimidinyl -NHCH2-Н 482. 4-pyrimidinyl -NHCH2-Н 483. 4-pyrimidinyl -NHCH2н 484. 4-pyrimidinyl -NHCH2-Н 485. 4-pyrimidinyl -NHCH2н 486. 4-pyrimidinyl -NHCH2-Н

Table 5. cont.

$$R^{2} \underbrace{\begin{array}{c} 0 \\ 5 \\ 4 \\ 3 \\ N \end{array}}_{N} R^{1}$$

5	#	R	Y	R ¹	R ²
	487.	4-pyrimidiny	l -NHCH ₂ -	CF ₂ CF ₃	Н
	488.	4-pyrimidiny	l -NHCH ₂ -	CF ₂ CF ₃	Н
	489.	4-pyrimidiny	l -NHCH ₂ -	CF ₂ CF ₃	Н
	490.	4-pyrimidiny	1 -NHCH ₂ -	H N	н
10	491.	N (CH	-NHCH ₂ -	CF ₂ CF ₃	Н
	492.	N (CH ₃)		CF ₂ CF ₃	н
	493.	$N^{(CH_3)}$		CF ₂ CF ₃	н
	494.	N (CH ₃)	-NHCH ₂ -	CF2CF3	Н
	495.	N N $(CH3)$	2 -NHCH ₂ -	$\bigcap_{0}^{CF_{2}CF_{3}}\bigcap_{N}$	Н

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Table 5. cont.

5				R	
Э	#	R	Y	R ¹	R ²
	496.	$N(CH_3)_2$	-NHCH ₂ -	CF ₂ CF ₃	Н
	497.	$N^{(CH_3)_2}$	-NHCH ₂ -	F ₃ C 0 N N CF ₂ CF ₃	Н
	498.	$\bigvee_{N}^{\text{CH}_3}$	-NHCH ₂ -	N N CH ₃	Н
10	499.	CH ₃	-NHCH ₂ -	N _{CH3}	Н
	500.	N	-NHCH ₂ -	CF ₂ CF ₃	Н
	501.	CH ₃	-NHCH ₂ -	N-CH3	Н
	502.	CH ₃	-NHCH ₂ -	CF ₂ CF ₃	Н
	503.	\mathcal{C}_{H^3}	-NHCH ₂ -	OH N	Н
15	504.	$\mathbb{C}^{\mathrm{H}_3}$	-NHCH ₂ -	F ₃ C CF ₃ N	Н

Table 5. cont.

$$\begin{array}{c|c} R^2 & O & N^{-R^1} \\ \hline \begin{pmatrix} 5 & 3 \\ 5 & 2 \end{pmatrix} & Y \\ 1 & 1 \\ \end{array}$$

5	#	R	Y	R ¹	R ²
	505.	SMe	-NHCH ₂ -	CF ₂ CF ₃	Н
	506.	SMe SMe	-NHCH ₂ -	CF ₂ CF ₃	Н
	507.	SMe	-NHCH ₂ -	ll N○	Н
	508.	r v	-NHCH ₂ -	N-CH ₃	H
10	509.	SMe	-NHCH ₂ -	CF ₂ CF ₃ N	н
	510.	SMe	-NHCH ₂ -	CF ₂ CF ₃	Н
	511.	OMe	-NHCH ₂ -	CF ₃	Н
	512.	OMe	-NHCH ₂ -	CF ₂ CF ₃	н
	513.	OMe	-NHCH ₂ -	CF ₂ CF ₃	Н

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Tabl 5. cont.

$$R^2 \underbrace{ \left(\begin{array}{c} 6 \\ 5 \\ 0 \end{array} \right)}_{N} R^1$$

5 R^2 R Y 514. -NHCH₂-Н 515. -NHCH₂-Н 516. -NHCH₂-Н 10 517. -NHCH₂-Н 518. -NHCH₂-Н -NHCH₂-519. Н 520. -NHCH₂-Н 521. -NHCH₂-Н 15 522. Н -NHCH₂-

Table 5. cont.

$$\mathbb{R}^2 \xrightarrow{\begin{array}{c} 0 \\ 5 \stackrel{4}{4} \end{array}} \mathbb{N}^{\mathbb{R}^1}$$

5	#	R	Y	R ¹	R ²
	523.	OMe	-NHCH ₂ -	CF ₂ CF ₃ H N	н
	524.	OMe	-NHCH2-	H ₃ C CH ₃	н
	525.	4-pyridyl	-NHCH ₂ -	2 OF mhomal	Н
	525.	4-bAtrdAT	-NACH2-	3-CF ₃ -phenyl	н
.0	526.	4-pyridyl	-NHCH ₂ -	CF,	Н
	527.	4-pyridyl	-NHCH ₂ -	CF ₃	Н
	528.	4-pyridyl	-NHCH ₂ -	CF3	Н
	529.	4-pyridyl	-NHCH₂-	CF ₃	Н
	530.	4-pyridyl	-NHCH ₂ -	O NH	Н

Table 5. cont.

$$R^2 \xrightarrow{\begin{pmatrix} 4 & 3 \\ 5 & 2 \end{pmatrix}} N^{R^1}$$

5 \mathbb{R}^2 Y 531. 4-pyridyl -NHCH₂-Н 532. 4-pyridyl -NHCH₂-Н 533. 4-pyridyl -NHCH₂-Н 534. 3-pyridyl -NH (CH₂)₂н 10 535. 3-pyridyl -NH(CH₂)₂-Н 536. 3-pyridyl -NH (CH₂)₂-

Н

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Table 5. cont.

5 \mathbb{R}^2 537. 3-pyridyl -NH (CH₂)₂-Н 538. 3-pyridyl 539. 4-pyridyl Н 540. 4-pyrimidinyl -NHCH2-541. 4-pyrimidinyl -NHCH2-Н 542. 4-pyrimidinyl -NHCH2-Н 543. 4-pyrimidinyl -NHCH2-Н

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Table 5. cont.

$$\begin{array}{c|c} R^2 & O & N \\ \hline & 5 & 3 \\ \hline & 5 & 2 \\ \hline & N & Y \\ \hline & R & \\ \end{array}$$

5	#	R	Y	R ¹	R ²
	544.	4-pyrimidinyl	-NHCH ₂ -	CF ₃	Н
	545.	4-pyrimidinyl	-NHCH ₂ -	CF ₃	Н
	546.	4-pyrimidinyl	-NHCH ₂ -	CF3	Н
	547.	4-pyrimidinyl	-NHCH ₂ -	N−CH ₃	Н
10	548.	4-pyrimidinyl	-NHCH ₂ -	N N CH3	Н
	549.	4-pyrimidinyl	-NHCH ₂ -	N _{CH3}	Н
	550.	U _N (3.3,72	-NHCH ₂ -	3-CF ₃ -phenyl	Н

557.

Table 5. cont.

-NHCH₂-

-NHCH₂-

3-CF₃-phenyl

Н

Table 5. cont.

$$R^{2} \underbrace{\begin{array}{c} 0 \\ 5 \\ 3 \\ 4 \\ 8 \\ N^{2} \end{array}}_{N} R^{1}$$

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Table 5. cont.

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Table 5. cont.

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Table 5. cont.

 R^1 \mathbb{R}^2 -NHCH₂-Н 578. -NHCH₂-Ħ 579. -NHCH₂-580. -NHCH₂-581. 4-pyrimidinyl -NHCH2-Н 582. 4-pyrimidinyl -NHCH2-Н

Н

583. 4-pyrimidinyl -NHCH2-

Table 5. cont.

-NHCH₂-

-NHCH₂-

Н

Н

10

588.

10

Table 5. cont.

Н

Table 5. cont.

$$\mathbb{R}^2 \underbrace{\begin{pmatrix} 5 & 3 \\ 5 & 2 \end{pmatrix}}_{N} \mathbb{R}^1$$

R1 Y 596. -NHCH₂-Н TOOTHOUT TOTHOOD н -NHCH₂-Н -NHCH₂-10 Н -NHCH₂-Н -NHCH₂-Н -NHCH₂-3-CF,-phenyl Н

-NHCH₂-

4-CF,-phenyl

Н

10

Table 5. cont.

$$\begin{array}{c|c}
R^2 & 0 & R^1 \\
\hline
5 & 3 & N & R^1 \\
\hline
N & 1 & R
\end{array}$$

#	R	Y	R ¹	R ²
604.	N NH NO	-NHCH ₂ -		Н
605.	NNH NH	-NHCH ₂ -	HN-N	Н
606.	NH NH NO	-NHCH ₂ -	FFF	Н
607.	NH: NH	-NHCH ₂ -	3-CF ₃ -phenyl	Н
608.	NH NH NO	-NHCH ₂ -	-	н
609.	N NH NO	-NHCH ₂ -	4-CF ₃ -phenyl	Н
610.		-NHCH ₂ -	N H	Н
611.		-NHCH ₂ -	HN-N	Н

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Tabl 5. cont.

$$R^2 \xrightarrow{\begin{cases} 4 & 3 \\ 5 & 2 \end{cases}} N^{R^1}$$

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			Ř	
#	R	Y	R ¹	R ²
612.	\smile	-NHCH ₂ -	3-CF,-phenyl	Н
	~~~~~			
613.	$\smile$	-NHCH ₂ -	4-CF3-phenyl	Н
614.	NH NH	-NHCH ₂ -	H ₃ C CH ₃	Н
615.	NH O NH	-NHCH ₂ -	-N	Н
616.	NH	-NHCH ₂ -	HN-N F	Н
617.	NH NH	-NHCH ₂ -	$\bigcap_{F} F$	Н
618.	NH	-NHCH ₂ -	3-CF,-phenyl	Н

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Table 5. cont.

			Ř	
#	R	Y	R ¹	R ²
619.	NH NH	-NHCH ₂ -	4-CF ₃ -phenyl	Н
620.	N O N CH3	-NHCH ₂ -	N H	Н
621.	CH ₃	-NHCH₂-	O-N	Н
622.	N O N CH3	-NHCH ₂ -	HN-N	Н
623.	CH ₃	-NHCH ₂ -	3-CF ₃ -phenyl	н
624.	CH ₃	-NHCH ₂ -	4-CF ₃ -phenyl	Н

Table 5. cont.

R² 4-pyridyl -NHCH₂-Н Н 627. н 10 628. Н 629. -NHCH₂-Н 630. -NHCH₂-Н

-NHCH₂-

н

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Table 5. cont.

$$R^{2} \xrightarrow{\begin{cases} 4 & 3 \\ 6 & 2 \end{cases}} N^{R^{1}}$$

# R Y R¹ R²

632. -NHCH₂- -NHCH₂- - H

634. 4-pyridyl -NHCH₂- - H

635. 4-pyridyl -NHCH₂- - H

include and the

Table 6.

	5	#	K-	n	R
		636.	4-chlorophenyl	1	6-I
		637.	3,4-dichlorophenyl	1	Н
		638.	4-fluorophenyl	1	Н
<u> </u> _		639.	3-chlorophenyl	1	Н
0	10	640.	3-fluorophenyl	1	H
.F		641.	3-fluoro-4-methoxyphenyl	1	H
(n En		642.	3-fluoro-4-methylphenyl	2	H
(0		643.	4-phenoxyphenyl	1	H
<b>ļ</b> ±		644.	3-phenoxyphenyl	1	H
g.	15	645.	4-biphenyl	1	Н
ļ-A.		646.	4-cyclohexylphenyl	1	H
<u>⊬</u> Ω		647.	2-quinolyl	1	H
0		648.	3-isoquinolyl	1	Н
ľV		649.	3-quinolyl	1	H
	20	650.	1-isoquinolyl	1	H
		651.	5-quinolyl	1	Н
		652.	5-isoquinolyl	1	H
		653.	6-quinolyl	1	H
		654.	6-isoquinolyl	1	Н
	25	655.	7-quinolyl	1	H
		656.	7-isoquinolyl	1	H
		657.	4-quinolyl	1	H

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Table 6. (cont.)

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5				
	#	R ¹	n	R ²
	658.	4-isoquinolyl	1	H
	659.	4-pyridyl	1	6-F
	660.	4-pyrimidinyl	1	H
10	661.	2-pyrimidinyl	1	Н
	662.	6-pyrimidinyl	1	Н
	663.	4-pyridazinyl	1	Н
	664.	5-pyridazinyl	1	H
	665.	4-indolyl	1	H
15	666.	5-isoindolyl	1	H
	667.	5-naphthyridinyl	1	Н
	668.	6-quinozalinyl	1	Н
	669.	6-isoquinolyl	1	Н
	670.	4-naphthyridinyl	1	Н
20	671.	5-quinozalinyl	1	Н
	672.	4-naphthyridinyl	1	H
	673.	tetrahydroquinolinyl	1	H
	674.	6-indazolyl	1	H
	675.	6-isoindolyl	1	Н
25	676.	5-indazolyl	1	H
	677.	5-isoindolyl	1	H
	678.	6-benzothienyl	1	H
	679.	6-benzofuryl	1	H
	680.	5-benzothienyl	1	H
30	681.	5-benzofuryl	1	H

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Table 6. (cont.)

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#	R ¹	n	R ²
682.	2-benzimidazolyl	1	Н
683.	2-benzoxazolyl	1	Н
684.	2-benzthiazolyl	1	Н
685.	6-benzimidazolyl	1	Н
686.	6-benzoxazolyl	1	Н
687.	6-benzthiazolyl	1	Н
688.	2-quinazolinyl	1	Н
689.	3-(phenoxy)-6-pyridyl	1	Н
690.	4-(phenylcarbonyl)phenyl	1	Н
691.	4-(phenylamino)phenyl	1	Н
692.	cyclohexyloxyphenyl	1	Н
693.	4-(3-thienyl)phenyl	1	Н
694.	4-(pyrazol-3-yl)phenyl	1	6-CH ₃

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5	#	R ¹	n	R ²
	695.	4-chlorophenyl	1	6-Cl
	696.	3,4-dichlorophenyl	1	5-C1
	697.	4-fluorophenyl	1	H
	698.	3-chlorophenyl	1	Н
10	699.	3-fluorophenyl	1	Н
	700.	3-fluoro-4-methoxyphenyl	1	H
	701.	3-fluoro-4-methylphenyl	1	H
	702.	4-phenoxyphenyl	1	H
	703.	3-phenoxyphenyl	1	Н
15	704.	4-biphenyl	1	Н
	705.	4-cyclohexylphenyl	1	Н
	706.	2-quinolyl	1	Н
	707.	3-isoquinolyl	1	H
	708.	3-quinolyl	1	H
20	709.	1-isoquinolyl	1	H
	710.	5-quinolyl	1	H
	711.	5-isoquinolyl	1	H
	712.	6-quinolyl	1	H
	713.	6-isoquinolyl	1	H
25	714.	7-quinolyl	1	H
	715.	7-isoquinolyl	1	H
	716.	4-quinolyl	1	Н
	717.	4-isoquinolyl	1	H

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Table 7. (cont.)

5	#	R ¹	n	R ²
	718.	4-pyridyl	1	Н
	719.	4-pyrimidinyl	1	H
	720.	2-pyrimidinyl	1	Н
	721.	6-pyrimidinyl	1	H
10	722.	4-pyridazinyl	1	H
	723.	5-pyridazinyl	1	Н
	724.	4-indolyl	1	Н
	725.	5-isoindolyl	1	Н
	726.	5-naphthyridinyl	1	Н
15	727.	6-quinozalinyl	1	Н
	728.	6-isoquinolyl	1	H
	729.	4-naphthyridinyl	1	Н
	730.	5-quinozalinyl	1	Н
	731.	4-naphthyridinyl	1	Н
20	732.	tetrahydroquinolinyl	1	Н
	733.	6-indazolyl	1	H
	734.	6-isoindolyl	1	H
	735.	5-indazolyl	1	Н
	736.	5-isoindolyl	1	Н
25	737.	6-benzothienyl	1	Н
	738.	6-benzofuryl	1	Н
	739.	5-benzothienyl	1	Н
	740.	5-benzofuryl	1	Н

# Table 8.

5	#	К		***
	741.	2-benzimidazolyl	1	Н
	742.	2-benzoxazolyl	1	Н
	743.	2-benzthiazolyl	1	Н
	744.	6-benzimidazolyl	1	Н
10	745.	6-benzoxazolyl	1	H
	746.	6-benzthiazoly16-benzoxazoly1	1	H
		2-quinazoliny16-benzoxazoly1	1	H
		3-(phenoxy)-6-pyridyl	1	H
		4-(phenylcarbonyl)phenyl	1	H
15		4-(phenylamino)phenyl	1	H
13		cyclohexyloxyphenyl	1	H
	752.		1	H
	753.		1	Н
	754.		1	${\tt EtO_2CCH=CH-}$
		4-chlorophenyl	1	5-Br
2.0	155.	4 - CITTOT Obugut 1		

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Table 8. (cont.)

5	#	R ¹	n	$\mathbb{R}^2$
	756.	4-pyridyl	1	Н
	757.	4-pyridyl	1	H
	758.	4-chlorophenyl	1	6-F
	759.	3,4-dichlorophenyl-	1	6-CH
10	760.	4-fluorophenyl	1	Н
	761.	3-chlorophenyl	1	H
	762.	3-fluorophenyl	1	H
	763.	3-fluoro-4-methoxyphenyl	1	Н
	764.	3-fluoro-4-methylphenyl	1	Н
15	765.	4-phenoxyphenyl	1	Н
	766.	3-phenoxyphenyl	1	H
	767.	4-biphenyl	1	Н
	768.	4-cyclohexylphenyl	1	Н
	769.	2-quinolyl	1	Н
20	770.	3-isoquinolyl	1	н
	771.	3-quinolyl	1	Н
	772.	1-isoquinolyl	1	Н
	773.	5-quinolyl	1	Н
	774.	5-isoquinolyl	1	Н
25	775.	6-quinolyl	1	Н
	776.	6-isoquinolyl	1	Н
	777.	7-quinolyl	1	Н
	778.	7-isoquinolyl	1	Н

Table 8. cont.

5  $\mathbb{R}^1$  $\mathbb{R}^2$ 779. 4-quinolyl Н 780. 4-isoquinolyl 1 Н 781. 4-pyridyl 1 Н 782. 4-pyrimidinyl 1 10 Н 783. 2-pyrimidinyl 1 н 784. 6-pyrimidinyl 1 Н 785. 4-pyridazinyl 1 Н 786. 5-pyridazinyl 1 н 787. 4-indolyl 15 788. 5-isoindolyl 1 789. 5-naphthyridinyl 790. 6-quinozalinyl 1 н 791. 6-isoquinolyl 1 20 792. 4-naphthyridinyl 793. 5-quinozalinyl 794. 4-naphthyridinyl 1 Н 795. 7-tetrahydroquinolinyl Н 796. 6-indazolyl 1 Н 25 797. 6-isoindolyl Н 798. 5-indazolyl 1 Н 799. 5-isoindolyl 1 Н 800. 6-benzothienyl 1 Н 801. 6-benzofuryl 1 Н

## Table 8. cont.

5	#	R ¹	n	R ²
	802.	5-benzothienyl	1	Н
	803.	5-benzofuryl	1	Н
	804.	2-benzimidazolyl	1	Н
	805.	2-benzoxazolyl	1	Н
10	806.	2-benzthiazolyl	1	Н
	807.	6-benzimidazolyl	1	Н
	808.	6-benzoxazolyl	1	Н
	809.	6-benzthiazolyl	1	Н
	810.	2-quinazolinyl	1	Н
15	811.	3-(phenoxy)-6-pyridyl	1	Н
	812.	4-(phenylcarbonyl)phenyl	1	Н
	813.	4-(phenylamino)phenyl	1	Н
	814.	4-cyclohexyloxyphenyl	1	Н
	815.	4-(3-thienyl)phenyl	1	Н
20	816.	4-(pyrazol-3-yl)phenyl	1	Н
	817.	3.4-dichlorophenvl	1	Н

# Table 9.

5	#	R ¹	n	R ²
	818.	4-chlorophenyl	1	6-F
	819.	3-fluoro-4-methoxyphenyl	1	H
	820.	4-phenoxyphenyl	1	Н
	821.	4-biphenyl	1	Н
10	822.	4-cyclohexylphenyl	1	Н
	823.	2-quinoly1	1	H
	824.	3-isoquinolyl	1	Н
	825.	3-quinolyl	1	Н

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# Example 826

# N-[4-(tert-Buty1)pheny1] (2-[(2,3-dihydrobenzo[b]furan-5-ylmethy1)amino](3-pyridy1))carboxamide

The titled compound was prepared from 2,3-dihydrobenzo[b]furan-5-ylmethylamine by the method described in Example 25. MS: (ES+) 402 (M+1)*; (ES-): 400 (M-1)⁻. Calc'd. for C₂₅H₂₇N₃O₂: 401.21.

### Example 827

{2-[(2,3-Dihydrobenzo[b]furan-5-ylmethyl)amino](3pyridyl)}-N-[3-(trifluoromethyl)phenyl]carboxamide

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The titled compound was prepared from 2,3-dihydrobenzo[b]furan-5-ylmethylamine by the method described in Example 25. MS: (ES+) 414 (M+1)*; (ES-): 412 (M-1)⁻. Calc'd. for C₂₂H₁₈F₁N₁O₂: 413.14.

The following compounds (Examples 828-864) were synthesized by the method described in Example 25 or Example 82 unless specifically described.

Table 10.

15	#	Y	R ¹	M+H	calc'd
	828.	-NH (CH ₂ ) ₂ -		375	374.2
	829.	-NH (CH ₂ ) ₂ -		375	374.2
	830.	-NH(CH ₂ ) ₂ -		411	410.2
	831.	-NH (CH ₂ ) ₂ -	FF	387	386.1
20	832.	-NH (CH ₂ ) ₂ -	<b>→</b>	361	360.2

Table 10. cont

5	#	Y	$R^1$	M+H	calc'd
	833.	-NH (CH ₂ ) ₂ -	CH ₃	457	456.6
	834.	-NH (CH ₂ ) ₂ -	−CF ₂ CF ₃	437	436.4
			H ₃ C _{CH₃}		
	835.	-NH (CH ₂ ) ₂ -	 сн _э	485.3	484.7
			H ₃ C CH ₃		
	836.	-NH (CH ₂ ) ₂ -	H H ₃ C _{CH₃}	388.3	387.5
10	837.	-NH (CH ₂ ) ₂ -	NH NH	485.3	484.6
			CF3		
	838.	-NH (CH ₂ ) ₂ -	NH	486	485.5

Table 10. cont.

5	#	Y	R ¹	M+H	calc'd
			CF₃		
			N-Boc		
	839.	-NH (CH ₂ ) ₂ -	CF ₂ CF ₃	586.4	585.6
	040	-NH (CH ₂ ) ₂ -		564	563.6
	040.	-NR (CR2/2-	CF ₂ CF ₃		
	0.41	-NH (CH ₂ ) ₂ -	OH N	580	579.6
	041.	=Nn (CH2/2-	CF2CF3		
	040	-NH (CH ₂ ) ₂ -	N _{CH₃}	564	563.6
	842.	-NH (CH ₂ ) ₂ -	H ₃ C CH ₃		
10	843.	-NH (CH ₂ ) ₂ -	N-CH ₃	499.2	498.7
			CF ₃		
	844	NH(CH ₂ ) ₂ -	N	512.1	511.6
	0		CF ₃		
			N-CH3		
	845	NH (CH ₂ ) ₂ -			497.6
	846	NHCH2-CH(4-mor	rpholino)		521.5
		-			

Table 10. cont.

M+H calc'd

5	# Y		
	847NH(CH ₂ ) ₂ -	CF ₃ N  CF ₂ CF ₃	514 513.6
	848NH(CH ₂ ) ₂ -	N-CH ₃	548.6
	849NH(CH ₂ ) ₂ -	H ₃ C CH ₃	484.1 483.2
	850NH(CH ₂ ) ₂ -	NH S=0	438 437
10	851NH(CH ₂ ) ₂ -	H ₃ C _C H ₃	430.2 429.5
	852NH(CH ₂ ) ₂ -	CF ₃	429 428.6
	853NH(CH ₂ ) ₂ -	N NH	498.5

Table 10. cont.

5	# Y	R ¹	M+H calc'd
	854NH (CH ₂ ) ₂ -	CF ₃	599 598.6
	855NH(CH ₂ ) ₂ -		471.3 470.7
	856NH(CH ₂ ) ₂ -		458 457.3
	857NH(CH ₂ ) ₂ -	N H ₃ C CH ₃	418.1 417.2
10	858NH(CH ₂ ) ₂ -		402 401.1
	859NH(CH ₂ ) ₂ -		445.9 445.5
	860NH(CH ₂ ) ₂ -	N H	432.1 431.

Table 10. cont.

#	Y	R ¹	M+H	Calc. u
		CF ₉	472 0	471.2
861.	-NH (CH ₂ ) ₂ -	H ₃ C CH ₃	1,2,,	
		NH NH		
862.	-NH (CH ₂ ) ₂ -	Ö	416.3	415.2
863	NH (CH ₂ ) ₂ -	CF3 OH	403.1	402.1
864	NH(CH ₂ ) ₂ -	F—VF	472.1	471.2

### Example 865

5 2-{[2-(1-Isopropyl-azetidin-3-ylmethoxy)-pyridin-4ylmethyl]-amino}-N-(4-trifluoromethyl-phenyl)-nicotinamide

A solution of 2-fluoro-N-(4-trifluoromethyl-phenyl)nicotinamide (107 mg) and [2-(1-isopropyl-azetidin-310 ylmethoxy)-pyridin-4-yl]-methylamine (89 mg) and NaHCO₃ (95
mg) was dissolved in IpOH (10 ml) and heated to 80°C for 18
h. After cooling to RT, the mixture was diluted with EtOAc
(50 ml) forming a precipitate which was filtered. The
filtrate was concentrated in vacuo. The residue was purified
15 by silica gel column chromatography (20% (12 N
NH₃/MeOH)/EtOAc) to give the product as a light yellow oil.
M+H 500.1: Calc'd 499.2.

The following compounds (Example 866-939) were synthesized 20 by the method described above.

866) N-(4-tert-Butyl-phenyl)-2-{{2-(1-isopropyl-azetidin-3ylmethoxy)-pyridin-4-ylmethyl}-amino}-nicotinamide.
M+H 488.1: Calc'd - 487.3

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- 867) 2-[(2,3-Dihydro-benzofuran-5-ylmethyl)-amino]-N-(4-[1-methyl-1-(1-methyl-piperidin-4-yl)-ethyl]-phenyl}nicotinamide. M+H 485.3: Calc'd 484.6.
- 868) N-(1-Acetyl-3,3-dimethyl-2,3-dihydro-1H-indol-6-yl)-2
  [(2,3-dihydro-benzofuran-5-ylmethyl)-amino]nicotinamide. M+H 457.1; Calc'd 456.5.
  - 869) 2-[(2,3-Dihydro-benzofuran-5-ylmethyl)-amino]-N-[3,3-dimethyl-1-(1-Boc-piperidin-4-ylmethyl)-2,3-dihydro-1H-indol-6-yl]-nicotinamide. M+H 612.6; Calc'd 611.8.
- 10 870) 2-[(2,3-Dihydro-benzofuran-5-ylmethyl)-amino]-N-[3,3-dimethyl-1-(1-methylpiperidin-4-ylmethyl)-2,3-dihydro-1H-indol-6-yl]-nicotinamide. M+H 526.3; Calc'd 525.7.
  - 871) N-(1-Acetyl-3,3-dimethyl-2,3-dihydro-1H-indol-6-yl)-2((2-[2-(1-methyl-piperidin-4-yl)-ethoxy]-pyridin-4ylmethyl)-amino)-nicotinamide. M+H Calc'd 556.
  - 872) 2-((2-[2-(1-Methyl-piperidin-4-yl)-ethoxy]-pyridin-4ylmethyl)-amino)-N-(3-trifluoromethyl-phenyl)nicotinamide. M+H Calc'd 513.
  - 873) N-(4-tert-Butyl-phenyl)-2-{[2-ethylpyridin-4-ylmethyl]amino}-nicotinamide.
  - 874) N-(4-tert-Butyl-phenyl)-2-((2-[2-(1-methyl-pyrrolidin-2-yl)-ethoxy]-pyridin-4-ylmethyl)-amino)-nicotinamide. M+H Calc'd 487.
  - 875) 2-((2-[2-(1-Methyl-pyrrolidin-2-yl)-ethoxy]-pyridin-4ylmethyl}-amino)-N-(4-pentafluoroethyl-phenyl)nicotinamide. M+H Calc'd 549.
  - 876) N-(4-Pentafluoroethyl-phenyl)-2-{[2-(2-pyrrolidin-1-yl-ethoxy)-pyridin-4-ylmethyl]-amino}-nicotinamide. M+H Calc'd 535.
- 30 877) N-(4-tert-Butyl-phenyl)-2-([2-(2-pyrrolidin-1-yl-ethoxy)-pyridin-4-ylmethyl]-amino)-nicotinamide. M+H Calc'd 473.

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- 878) N-[3-(4-Boc-piperazin-1-ylmethyl)-5-trifluoromethyl-phenyl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide.

  M+H 571 4: Calc'd 570 3
- 879) N-[3-(4-Boc-piperazine-1-carbonyl)-5-trifluoromethylphenyl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide.
  M+H Calc'd 584.
  - 880) N-[3-(4-Boc-piperazine-1-carbonyl)-5-trifluoromethyl-phenyl]-2-(2-pyridin-4-yl-ethylamino)-nicotinamide.
    M+H Calc'd 598.
- 10 881) N-[3-(4-Methyl-piperazin-1-ylmethyl)-4pentafluoroethyl-phenyl]-2-[(pyridin-4-ylmethyl)aminol-nicotinamide. M+H Calc'd 534.
  - 882) N-[3-(4-Boc-piperazin-1-ylmethyl)-4-pentafluoroethyl-phenyl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide.
    M+H 621.4: Calc'd 620.
  - 883) 2-{[2-(1-Methyl-piperidin-4-ylmethoxy)-pyridin-4-ylmethyl]-amino}-N-(4-trifluoromethyl-phenyl)-nicotinamide.
  - 884) N-(4-tert-Butyl-phenyl)-2-{[2-(1-methyl-piperidin-4-ylmethoxy)-pyridin-4-ylmethyl]-amino}-nicotinamide.
  - 885) 2-((2-[3-(1-Methyl-piperidin-4-yl)-propoxy]-pyridin-4ylmethyl)-amino)-N-(4-pentafluoroethyl-phenyl)nicotinamide. M+H 578.3. Calc'd 577.2.
  - 886) N-(1-Acetyl-3,3-dimethyl-2,3-dihydro-1H-indol-6-yl)-2[(2-methoxy-pyridin-4-vlmethyl)-aminol-nicotinamide.
  - 887) N-[3,3-Dimethyl-1-(1-methyl-piperidin-4-yl)-2,3-dihydro-1H-indol-6-yl]-2-[(2-methoxy-pyridin-4-ylmethyl)-amino]-nicotinamide. M+H 501.2; Calc'd 500.3.
- 30 888) N-(1-Boc-3,3-dimethyl-2,3-dihydro-1H-indol-6-yl)-2-[(2-methoxy-pyridin-4-ylmethyl)-aminol-nicotinamide.
  - 889) N-[3,3-Dimethyl-1-(1-Boc-piperidin-4-ylmethyl)-2,3-dihvdro-1H-indol-6-yl]-2-[(2-methoxy-pyridin-4-

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ylmethyl)-amino]-nicotinamide. M+H 601.6; Calc'd 600.34

- 890) N-[3,3-Dimethyl-1-(1-methyl-piperidin-4-yl)-2,3dihydro-1H-indol-6-yl]-2-[(pyridin-4-ylmethyl)-amino]nicotinamide.
- 891) N-[1-(2-Dimethylamino-acetyl)-3,3-dimethyl-2,3-dihydro-1H-indol-6-yl]-2-[(2-methoxy-pyridin-4-ylmethyl)aminol-nicotinamide.
- 892) N-[1-(2-Dimethylamino-acetyl)-3,3-dimethyl-2,3-dihydro10 1H-indol-6-yl]-2-[(pyridin-4-ylmethyl)-amino]nicotinamide.
  - 893) 2-[(2-Methoxy-pyridin-4-ylmethyl)-amino]-N-[3-(1-Bocpiperidin-4-ylmethoxy)-5-trifluoromethyl-phenyl]nicotinamide
- 15 894) N-[3,3-Dimethyl-1-(1-Boc-pyrrolidin-2-ylmethoxy)-2,3-dihydro-1H-indol-6-yl]-2-[(2-methoxy-pyridin-4-ylmethyl)-amino]-nicotinamide.
  - 895) N-[3,3-Dimethyl-1-(2-Boc-amino-acetyl)-2,3-dihydro-1H-indol-6-yl]-2-[(2-methoxy-pyridin-4-ylmethyl)-amino]-nicotinamide.
  - 896) N-[3,3-Dimethyl-1-(2-Boc-amino-acetyl)-2,3-dihydro-1Hindol-6-yl]-2-[(pyridin-4-ylmethyl)-amino]nicotinamide.
  - 897) 2-[(2-Methoxy-pyridin-4-ylmethyl)-amino]-N-[3-(1methyl-pyrrolidin-2-ylmethoxy)-5-trifluoromethylphenyl]-nicotinamide. M+H 516.1.
  - 898) 2-[(2-Methoxy-pyridin-4-ylmethyl)-amino]-N-[3-(1-Bocpiperidin-4-ylmethyl)-5-trifluoromethyl-phenyl]nicotinamide. M+H 501.3.
- 30 899) 2-[(2-Methoxy-pyridin-4-ylmethyl)-amino]-N-[3-(4-Bocpiperazin-1-ylmethyl)-5-trifluoromethyl-phenyl]nicotinamide.

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900) 2-{[2-(3-Morpholin-4-vl-propoxy)-pyridin-4-vlmethvl]amino)-N-(4-pentafluoroethyl-phenyl)-nicotinamide, M+H 566

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- 901) (S) 2-{[2-(1-Methyl-pyrrolidin-2-ylmethoxy)-pyridin-4-5 vlmethvll-amino}-N-(4-pentafluoroethvl-phenvl)nicotinamide, M+H 536.
  - 902) N-(3-tert-Butyl-isoxazol-5-yl)-2-{[2-(3-morpholin-4-ylpropoxy)-pyridin-4-vlmethyll-amino}-nicotinamide. M+H 495. Calc'd 494.
- 10 903) N-(1-Acetyl-3,3-dimethyl-2,3-dihydro-1H-indol-6-yl)-2-{[2-(3-morpholin-4-vl-propylamino)-pyridin-4vlmethyl]-amino}-nicotinamide. M+H 558; Calc'd 557.
  - 904) N-(4-tert-Butvl-phenvl)-2-{[2-(3-morpholin-4-vlpropoxy)-pyridin-4-ylmethyl]-amino}-nicotinamide. 504. Calc'd 503.
  - 905) N-(4-tert-Butvl-phenvl)-2-([2-(2-morpholin-4-vlethoxy)-pyridin-4-ylmethyl]-amino}-nicotinamide. M+H 409: Calc'd 489.
- 906) 2-{[2-(2-Morpholin-4-vl-ethoxy)-pyridin-4-vlmethyl]-20 amino}-N-(4-trifluoromethyl-phenyl)-nicotinamide. M+H 502 · Calc'd 501
  - 907) 2-{[2-(2-Morpholin-4-yl-ethoxy)-pyridin-4-ylmethyl]amino}-N-(3-trifluoromethyl-phenyl)-nicotinamide. M+H 502: Calc'd 501.
- 25 908) 2-{[2-(2-Morpholin-4-yl-ethoxy)-pyridin-4-ylmethyl]amino}-N-(4-pentafluoroethyl-phenyl)-nicotinamide. M+H 552; Calc'd 551.
  - 909) N-(3-tert-Butvl-isoxazol-5-vl)-2-{[2-(2-morpholin-4-vlethoxy)-pyridin-4-ylmethyl]-amino}-nicotinamide. M+H 481: Calc'd 480.
  - 910) N-(1-Acetyl-3,3-dimethyl-2,3-dihydro-1H-indol-6-yl)-2-{[2-(2-morpholin-4-yl-ethoxy)-pyridin-4-ylmethyl]amino}-nicotinamide, M+H 545; Calc'd 544.

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- 911) N-(1-Acetyl-3,3-dimethyl-2,3-dihydro-1H-indol-6-yl)-2{[2-(1-methyl-piperidin-4-yloxy)-pyridin-4-ylmethyl]amino)-nicotinamide.
- 912) 2-{[2-(1-Methyl-piperidin-4-yloxy)-pyridin-4-ylmethyl]amino}-N-(4-trifluoromethyl-phenyl)-nicotinamide.
- 913) 2-{[2-(1-Methyl-piperidin-4-yloxy)-pyridin-4-ylmethyl]amino}-N-(4-pentafluoroethyl-phenyl)-nicotinamide.
- 914) 2-([2-(1-Methyl-piperidin-4-yloxy)-pyridin-4-ylmethyl]amino)-N-(4-tert-butvl-phenvl)-nicotinamide.
- 10 915)(R) N-(4-tert-Butyl-phenyl)-2-([2-(1-methyl-pyrrolidin-2-ylmethoxy)-pyridin-4-ylmethyl]-amino)-nicotinamide. M+H 474: Calc'd 473.
  - 916) (R) N-[3-(1-Boc-pyrrolidin-2-ylmethoxy)-5trifluoromethyl-phenyl]-2-[(pyridin-4-ylmethyl)aminol-nicotinamide.
  - 917) (R) N-[3-(1-Methyl-pyrrolidin-2-ylmethoxy)-5trifluoromethyl-phenyl]-2-[(pyridin-4-ylmethyl)amino]-nicotinamide. M+H 486; Calc'd 485.5.
  - 918) N-[3-(1-Methyl-piperidin-4-yloxy)-5-trifluoromethyl-phenyl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide.
  - 919) N-[3-(1-Methyl-piperidin-4-ylmethyl)-5-trifluoromethyl-phenyl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide.
- 920) N-[4-tert-Butyl-3-(1-Boc-pyrrolidin-2-ylmethoxy)phenyl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide.

  25 M+H 560; Calc'd 559.
  - 921) N-(3,3-Dimethyl-2,3-dihydro-benzofuran-6-yl)-2-{[2-(1-methyl-piperidin-4-ylmethoxy)-pyridin-4-ylmethyl]-amino}-nicotinamide.
- 922) 2-((2-[3-(1-Methyl-piperidin-4-yl)-propoxy]-pyridin-4-30 ylmethyl)-amino)-N-(4-trifluoromethyl-phenyl)nicotinamide.
  - 923) 2-((2-[3-(1-Methyl-piperidin-4-yl)-propoxy]-pyridin-4-ylmethyl)-amino)-N-(3-trifluoromethyl-phenyl)nicotinamide.

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- 924) 2-((2-[3-(1-Methyl-piperidin-4-yl)-propoxy]-pyridin-4-ylmethyl}-amino)-N-(4-tert-butyl-phenyl)-nicotinamide.
- 925) 2-({2-[3-(1-Methyl-piperidin-4-yl)-propoxy}-pyridin-4ylmethyl}-amino)-N-(3-tert-butyl-isoxazol-5-yl)nicotinamide.
- 926) N-(3,3-Dimethyl-2,3-dihydro-1H-indol-6-yl)-2-({2-[3-(1-methyl-piperidin-4-yl)-propoxy]-pyridin-4-ylmethyl}-aminol-nicotinamide.
- 927) 2-[(Pyridin-4-ylmethyl)-amino]-N-(3,9,9-trimethyl10 2,3,4,4a,9,9a-hexahydro-1H-3-aza-fluoren-6-yl)nicotinamide.
  - 928) N-[3,3-Dimethyl-1-(1-Boc-piperidin-4-ylmethyl)-2,3dihydro-1H-indol-6-yl]-2-[(pyridin-4-ylmethyl)-amino]nicotinamide
- 929) N-[3,3-Dimethyl-1-(1-methyl-piperidin-4-ylmethyl)-2,3-dihydro-1H-indol-6-yl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide. M+H 485.3; Calc'd 484.6.
- 20 by the method described above, substituting K₂CO₃ for NaHCO₃.
  930) 2-{[2-(1-Methyl-piperidin-4-ylmethoxy)-pyridin-4-ylmethyl]-amino}-N-(4-pentafluoroethyl-phenyl)-

The following compounds (Example 930-937) were synthesized

- 25 932) N-(1-Acetyl-3,3-dimethyl-2,3-dihydro-1H-indol-6-yl)-2-([2-(1-methyl-piperidin-4-ylmethoxy)-pyridin-4ylmethyl]-amino)-nicotinamide. M+H 543.4; Calc'd 542.3.
- 933) N-(4-tert-Butyl-phenyl)-2-([2-(3-morpholin-4-yl-30 propylamino)-pyrimidin-4-ylmethyl]-amino)nicotinamide. M+H 504.3: Calc'd 503.6.

nicotinamide, M+H 550.2: Calc'd 549.2.

934) 2-([2-(3-Morpholin-4-yl-propylamino)-pyrimidin-4-ylmethyl]-amino)-N-(4-pentafluoroethyl-phenyl)-nicotinamide. M+H 566.3: Calc'd 565.55.

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935) 2-{[2-(3-Morpholin-4-yl-propylamino)-pyrimidin-4-ylmethyl]-amino)-N-(3-trifluoromethyl-phenyl)-nicotinamide. M+H 516.0: Calc'd 515.5.

- 936) N-(4-tert-Butyl-phenyl)-2-({2-[2-(1-methyl-pyrrolidin-2-yl)-ethylamino]-pyrimidin-4-ylmethyl)-amino)nicotinamide, M+H Calc'd 487.6.
- 937) N-(1-Acetyl-3,3-dimethyl-2,3-dihydro-1H-indol-6-yl)-2-((2-[2-(1-methyl-pyrrolidin-2-yl)-ethylamino]pyrimidin-4-ylmethyl)-amino)-nicotinamide. M+H Calc'd 542.69.

The following compounds (Example 938-939) were synthesized by the method described above, substituting  $Cs_2CO_3$  for NaHCO1.

938) 2-((2-(1-Methyl-piperidin-4-ylmethoxy)-pyridin-4-ylmethyl]-amino}-N-[3-(1-methyl-piperidin-4-yl)-5-trifluoromethyl-phenyl]-nicotinamide. M H 597.0;

20 939) N-(3-tert-Butyl-isoxazol-5-yl)-2-{[2-(1-methyl-piperidin-4-ylmethoxy)-pyridin-4-ylmethyl]-amino}-nicotinamide. M+H 479: Calc'd 478.3..

Calc'd 596.7.

The following compounds (Example 940-945) were synthesized by the method described above, substituting t-BuOH for IpOH.

- 940) N-[3-(1-Boc-azetidin-3-ylmethoxy)-5-trifluoromethyl-phenyl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide. M+H 558.1. Calc'd 557.6.
- 30 941) 2-[(2-Methoxy-pyridin-4-ylmethyl)-amino]-N-[3-(1-Boc-azetidin-3-ylmethoxy)-5-trifluoromethyl-phenyl]-nicotinamide. M+H 588.1. Calc'd 587.2.

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- 942) 2-[(Pyridin-4-ylmethyl)-amino]-N-(2,2,4-trimethyl-3,4-dihydro-2H-benzo[1,4]oxazin-6-yl)-nicotinamide. M+H
- 943) N-(4-Acetyl-2,2-dimethyl-3,4-dihydro-2H-
- 5 benzo[1,4]oxazin-6-yl)-2-[(pyridin-4-ylmethyl)-amino]nicotinamide. M+H 432.1; Calc'd 431.5.
  - 944) N-(2,2-Dimethyl-3-oxo-3,4-dihydro-2H-benzo[1,4]oxazin-6-yl)-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide. M+H 404.5: Calc'd 403.2.
- 10 945) 2-[[2-(1-Benzhydryl-azetidin-3-yloxy)-pyridin-4-ylmethyl]-amino)-N-(4-tert-butyl-phenyl)-nicotinamide.

  M+H 598 4 Calc/d 597 3

The following compounds (Example 946-993) were synthesized by the method described above, unless specifically described.

- 946) N-(4,4-Dimethyl-1-oxo-1,2,3,4-tetrahydro-isoquinolin-7-yl)-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide was prepared with MeOH as the solvent at 110°C. M+H 402.3.
- 947) N-(4-tert-Butyl-phenyl)-2-((2-[2-(1-methyl-piperidin-4-yl)-ethoxyl-pyridin-4-ylmethyl)-amino)-nicotinamide was prepared with pentanol at 95°C. M+H Calc'd 501.
- 948) N-(3-tert-Butyl-isoxazol-5-yl)-2-((2-[2-(1-methyl-piperidin-4-yl)-ethoxy]-pyridin-4-ylmethyl)-amino)-nicotinamide was prepared with pyridine at 95°C. M+H Calc'd 492.
- 949) N-(3-trifluoromethylphenyl)-2-((2-[2-(1-methyl-piperidin-4-yl)-ethoxy]-pyridin-4-ylmethyl)-amino)nicotinamide was prepared with pyridine at 95°C. M+H
- 950) 2-[(2,3-Dihydro-benzofuran-6-ylmethyl)-amino]-N-[3-(1-Boc-pyrrolidin-2-ylmethoxy)-4-pentafluoroethyl-

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phenyl]-nicotinamide was prepared with DIEA at 120°C.
M+H 663.4: Calc'd 662.6.

- 951) (R) N-[3-(2-Hydroxy-3-pyrrolidin-1-yl-propoxy)-4-pentafluoroethyl-phenyl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide was prepared with IpOH as the solvent at 135°C M+H 566 5: Calc'd 565.5.
- 952) (S) N-[3-(2-Hydroxy-3-pyrrolidin-1-yl-propoxy)-4-pentafluoroethyl-phenyl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide was prepared with IpOH as the solvent at 135°C. M+H 566.5; Calc'd 565.5.
- 953) N-[4-tert-Butyl-3-(1-methyl-piperidin-4-ylmethoxy)phenyl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide
  was prepared with IpOH as the solvent at 130°C. M+H
  468.3; Calc'd 487.6.
- 954) N-[3-(1-Methyl-piperidin-4-ylmethoxy)-4pentafluoroethyl-phenyl]-2-[(pyridin-4-ylmethyl)amino]-nicotinamide was prepared with IpOH as the solvent at 135°C. M+H 550.2; Calc'd 549.5.
  - 955) N-[4-Pentafluoroethyl-3-(2-piperidin-1-yl-ethoxy)phenyl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide
    was prepared with IpOH as the solvent at 130°C. M+H
    550.1; Calc'd 549.5.
  - 956) N-[4-Trifluoromethyl-3-(2-piperidin-1-yl-ethoxy)-phenyl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide was prepared with IpOH as the solvent at 130°C. M+H 486.3; Calc'd 485.5.
    - 957) (S) N-[3-(1-Boc-pyrrolidin-2-ylmethoxy)-4pentafluoroethyl-phenyl]-2-[(pyridin-4-ylmethyl)amino]-nicotinamide was prepared with DIEA at 135°C.
      M+H 572. Calc'd 571.6.
    - 958) (R) N-[3-(1-Boc-pyrrolidin-2-ylmethoxy)-4trifluoromethyl-phenyl]-2-[(pyridin-4-ylmethyl)amino]-nicotinamide was prepared with DIEA at 130°C. M+H 622. Calc'd 621.6.

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- 959) (R) N-[3-(1-Boc-pyrrolidin-2-ylmethoxy)-4-pentafluoroethyl-phenyl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide was prepared with DIEA at 130°C.

  M+H 622.4. Calc'd 621.6.
- 5 960) N-(4-tert-Butyl-phenyl)-2-{[2-(1-methyl-piperidin-4-yloxy)-pyridin-4-ylmethyl]-amino}-nicotinamide was prepared with pyridine and TEA at 90°C.M+H 474.
  - 961) N-(3-Trifluoromethyl-phenyl)-2-([2-(1-methyl-piperidin-4-yloxy)-pyridin-4-ylmethyl]-amino)-nicotinamide was prepared with pyridine and TEA at 90°C. M+H 486.
  - 962) N-(3-tert-Butyl-isoxazol-5-yl)-2-{[2-(1-methyl-piperidin-4-yloxy)-pyridin-4-ylmethyl]-amino}nicotinamide was prepared with pyridine and TEA at
    90°C. M+H 465.
- 963) N-[3-(3-Piperidin-1-yl-propy1)-5-trifluoromethyl-phenyl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide was prepared with pyridine at 90°C. M+H 498; Calc'd 497.6.
- 964) N-[3-(3-Morpholin-4-y1-propy1)-5-trifluoromethy120 pheny1]-2-[(pyridin-4-y1methy1)-amino]-nicotinamide
  was prepared at 130°C neat. M+H 500. Calc'd 499.2.
  - 965) 2-[(2-Methoxy-pyridin-4-ylmethyl)-amino]-N-[3-(1-Boc-piperidin-4-yloxy)-5-trifluoromethyl-phenyl]-nicotinamide was prepared at 130°C neat. M+H 602. Calc'd for CnHuFnNos: 601.6.
  - 967) N-{4-tert-Butyl-3-[2-(1-Boc-piperidin-4-yl)-ethoxy]-phenyl}-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide was prepared with DIEA and IDOH at 130°C. M+H 574.6.
- 30 968) N-[4-tert-Butyl-3-(1-methyl-azetidin-3-ylmethoxy)-phenyl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide was prepared with IpOH and DIEA at 130°C. M+H 546.
  - 969) N-(3,3-Dimethyl-1,1-dioxo-2,3-dihydro-1H-1λ-benzo[d]isothiazol-6-yl)-2-[(pyridin-4-ylmethyl)-

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- amino]-nicotinamide was prepared neat at 130°C. M+H 424: Calc'd 423.
- 970) N-[1,1,4,4-Tetramethyl-1,2,3,4-tetrahydro-naphth-6-yl]2-[(pyridin-4-ylmethyl)-amino]-nicotinamide was
  prepared neat at 130°C. M+H 415; Calc'd 414.
- 971) N-{4-[1-Methyl-1-(1-methyl-piperidin-4-yl)-ethyl]phenyl}-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide
  was prepared with pyridine. M+H 444; Calc'd 443.27.
- 972) 2-[(2-Methoxy-pyridin-4-ylmethyl)-amino]-N-{4-[1-10 methyl-1-(1-methyl-piperidin-4-yl)-ethyl]-phenyl)nicotinamide was prepared with pyridine and NaHCO₃ at 110°C. MS: 473 (M+H). Calc'd for C₂₂H₃₅N₅O₂ - 472.6.
  - 973) N-(3,3-Dimethyl-2,3-dihydro-benzofuran-6-yl)-2[(pyridin-4-ylmethyl)-amino]-nicotinamide was prepared
    with IbOH at 120°C. M+H 375. Calc'd for C>+H+NAO: 374.
  - 974) 2-{[2-(3-Dimethylamino-propoxy)-pyridin-4-ylmethyl]-amino)-N-(4-pentafluoroethyl-phenyl)-nicotinamide. M+H 524; Calc'd 523.2.
  - 975) N-[3-(1-Methyl-piperidin-4-yl)-5-trifluoromethylphenyl]-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide. M+H 470.4: Calc'd 469.21.
    - 976) 2-{[2-(1-Methyl-piperidin-4-ylmethoxy)-pyridin-4-ylmethyl]-amino}-N-[3-(1-methyl-piperidin-4-yl)-5-trifluoromethyl-phenyl]-nicotinamide. M+H 597.0;
      Calc'd 596.31.
    - 977) N-[3-(azetidin-3-ylmethoxy)-5-trifluoromethyl-phenyl]2-[(pyridin-4-ylmethyl)-amino]-nicotinamide. M+H
      458.1: Calc'd 457.2.
- 978) N-(3-Hydroxy-5-trifluoromethyl-phenyl)-2-[(pyridin-4-30 ylmethyl)-amino]-nicotinamide. M+H 388.9; Calc'd 388.11.
  - 979) N-(2-Acetyl-4,4-dimethyl-1,2,3,4-tetrahydroisoquinolin-7-yl)-2-[(pyridin-4-ylmethyl)-amino]nicotinamide. M+H 430; Calc'd 429.22.

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- 980) N-[2-(4-methoxy-benzyl)-4,4-dimethyl-1-oxo-1,2,3,4-tetrahydro-isoquinolin-7-yl]-2-[(pyridin-4-ylmethyl)-aminol-nicotinamide. M+H 522.3: Calc'd 521.24.
- 981) N-(2-Acetyl-4,4-dimethyl-1,2,3,4-tetrahydroisoquinolin-7-yl)-2-[(quinolin-4-ylmethyl)-amino]benzamide. M+H 479: Calc'd 478.24.
  - 982) 2-[(Pyridin-4-ylmethyl)-amino]-N-[3-(2-pyrrolidin-1-ylethoxy)-4-trifluoromethyl-phenyl]-nicotinamide. M+H 486: Calc'd 485.
- 10 983) 2-{[2-(1-Methyl-piperidin-4-ylmethoxy)-pyridin-4ylmethyl]-amino)-N-(3-trifluoromethyl-phenyl)nicotinamide. M+H 500.5; Calc'd 499.5.
  - 984) N-[3-(1-Boc-azetidin-3-ylmethoxy)-4-tert-butyl-phenyl]2-[(pyridin-4-ylmethyl)-amino]-nicotinamide. M+H 546.
    Calc'd 545.
  - 985) 2-Methyl-2-[4-((2-[(pyridin-4-ylmethyl)-amino]pyridine-3-carbonyl)-amino)-phenyl]-propionic acid methyl ester. M+H 405; Calc'd 404.
- 986) N-(4-tert-Butyl-phenyl)-2-{[2-(3-morpholin-4-yl-20 propylamino)-pyrimidin-4-ylmethyl]-amino}nicotinamide. M+H 504.3; Calc'd 503.
  - 987) N-(4-pentafluoroethyl-phenyl)-2-{[2-(3-morpholin-4-yl-propylamino)-pyrimidin-4-ylmethyl]-amino}-nicotinamide. M+H 566.3; Calc'd 565.
- - 989) N-(4-tert-Butyl-phenyl)-2-((2-[2-(1-methyl-pyrrolidin-2-yl)-ethylamino]-pyrimidin-4-ylmethyl)-amino)nicotinamide. M+H 488.4; Calc'd 487.
  - 990) N-(1-Acetyl-3,3-dimethyl-2,3-dihydro-1H-indol-6-yl)-2-((2-[2-(1-methyl-pyrrolidin-2-yl)-ethylamino]pyrimidin-4-ylmethyl)-amino)-nicotinamide. M+H 543.5; Calc'd 542.

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991) N-(1-Acetyl-3,3-dimethyl-2,3-dihydro-1H-indol-6-yl)-2-(2,3-dihydro-benzo[1,4]dioxin-6-ylamino)-nicotinamide. M+H 459.3.

992) 2-({2-[2-(1-Methyl-pyrrolidin-2-yl)-ethylamino}pyrimidin-4-ylmethyl)-amino)-N-(3-trifluoromethylphenyl)-nicotinamide. M+H 500.4; Calc'd 499.

### Example 993

N-(3,3-Dimethy1-2,3-dihydro-1H-indo1-6-y1)-2-({2-[2-(1-methy1-piperidin-4-y1)-ethoxy]-pyridin-4-ylmethy1}amino)-nicotinamide

N-(1-Acetyl-3,3-dimethyl-2,3-dihydro-1H-indol-6-yl)-2-((2-[2-(1-methyl-piperidin-4-yl)-ethoxy]-pyridin-4-ylmethyl)-amino)-nicotinamide (300 mg, Example 871) was dissolved in conc. HCl (20 ml) and EtOH (20 ML) and heated at  $70^{\circ}\text{C}$  for 4 H. The mixture was concentrated and the residue was diluted with sat'd NaHCO₃ and CH₂Cl₂. The organic layer was dried over Na₂SO₄ and concentrated to obtain the desired compound. M+H 515, Calc'd for C₃H₃₈N₆O₂: 514.

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The following compounds (Example 995-1009) were synthesized by the method described above, unless specifically described.

- 5 995) N-(2,2-Dimethyl-3,4-dihydro-2H-benzo[1,4]oxazin-6-yl)2-[(pyridin-4-ylmethyl)-amino]-nicotinamide. M+H
  390.3: Calc'd 389.4.
  - 996) N-(4,4-Dimethyl-1,2,3,4-tetrahydro-isoquinolin-7-yl)-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide. M+H 388.3.
- 10 997) N-(3,3-Dimethyl-2,3-dihydro-1H-indol-6-yl)-2-{[2-(1-methyl-piperidin-4-ylmethoxy)-pyridin-4-ylmethyl]-amino}-nicotinamide. M+H 501.3: Calc'd 500.3.
  - 998) N-(3,3-Dimethyl-1-piperidin-4-yl-2,3-dihydro-1H-indol-6-yl)-2-[(pyridin-4-ylmethyl)-amino]-nicotinamide was prepared with 1N HCl in ether and dioxane at RT. M+H 457.2: Calc'd 456.7.
  - 999) N-(3,3-dimethy1-2,3-dihydro-1H-indo1-6-y1)-2-({2-[2-(1-methy1-pyrrolidin-2-y1)-ethylamino]-pyrimidin-4-ylmethy1}-amino)-nicotinamide. M+H Calc'd 500.65.
- 20 1000) N-(3,3-dimethyl-2,3-dihydro-1H-indol-6-yl)-2-[(2-methoxy-pyridin-4-ylmethyl)-amino]-nicotinamide. M+H 404.3: Calc'd 403.2.
  - 1001) N-[3,3-Dimethyl-1-(piperidin-4-ylmethyl)-2,3-dihydro-1H-indol-6-yl]-2-[(2-methoxy-pyridin-4-ylmethyl)amino]-nicotinamide was prepared with HCl in EtOAc. M+H 501.4; Calc'd 500.3.
  - 1002) N-(3,3-Dimethyl-1-piperidin-4-yl-2,3-dihydro-1H-indol-6-yl)-2-[(2-methoxy-pyridin-4-ylmethyl)-amino]nicotinamide. M+H 487.4: Calc'd 486.3.
- 30 1003) 2-[(2-Methoxy-pyridin-4-ylmethyl)-amino]-N-[3-(piperidin-4-ylmethoxy)-5-trifluoromethyl-phenyl)nicotinamide was prepared with HCl in EtOAc.

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- 1004) N-[3,3-Dimethyl-1-(pyrrolidin-2-ylmethyl)-2,3-dihydro-1H-indol-6-yl]-2-[(2-methoxy-pyridin-4-ylmethyl)aminol-nicotinamide was prepared with HCl in EtOAc.
- 1006) N-(3,3-dimethyl-2,3-dihydro-1H-indol-6-yl)-2-{[2-(2morpholin-4-yl-ethoxy)-pyridin-4-ylmethyl]-amino}nicotinamide. M+H 503; Calc'd 502.
- 1007) N-(3,3-dimethyl-2,3-dihydro-1H-indol-6-yl)-2-{[2-(1-methyl-piperidin-4-yloxy)-pyridin-4-ylmethyl}-amino)nicotinamide. M+H 529.
- 1008) N-(3,3-dimethyl-2,3-dihydro-1H-indol-6-yl)-2-{[2-(2-morpholin-4-yl-propylamino)-pyridin-4-ylmethyl]amino)-nicotinamide. M+H 516; Calc'd 515.
  - 1009) N-(3,3-dimethyl-2,3-dihydro-1H-indol-6-yl)-2-({2-[2-(1-methyl-pyrrolidin-2-yl)-ethylamino]-pyrimidin-4ylmethyl}-amino)-nicotinamide. M+H 501.4; Calc'd 500.

Example 1010

N-(4-Pentafluoroethyl-phenyl)-2-[(pyrimidin-4-ylmethyl)amino]-nicotinamide

2-Amino-N-(4-pentafluoroethyl-phenyl)-nicotinamide (180 mg), TsOH (40 mg) and a solution of pyrimidine-4-carboxaldehyde in DMSO (10 ml) were stirred at 60 C for 6 h. Treated with NaBH₄ (200 mg) and stirred for 2 h at RT. MS (ES * ): 566.3 (M+H) * ; Calc'd for  $C_{2k}H_{2k}F_kN_{2k}N_{2k}$  - 565.

### Example 1011

2-{[2-(Azetidin-3-yloxy)-pyridin-4-ylmethyl]-amino}-N-(4tert-butyl-phenyl)nicotinamide

2-{[2-(1-Benzhydry1-azetidin-3-yloxy)-pyridin-415 ylmethyl]-amino}-N-(4-tert-buty1-phenyl)-nicotinamide (210
mg) was heated at reflux with Et,SiH (5 ml) and TFA (15 ml)
for 9 h. The mixture was concentrated, then diluted with
CH₂Cl₂ (50 ml) and washed with sat'd NaHCO₃ (50 ml) and brine
(30 ml), dried over MgSO₄ and purified by silica gel
20 chromatography (10% MeOH/2M NH₃ 90% EtOAc) to afford the
product as a yellow solid. M+H Calc'd for C₂₅H₂₉N₅O₂: 431.2.

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### Example 1012

N-(2,3,3-Trimethyl-1,1-dioxo-2,3-dihydro-1H-1\lambda'benzo[d]isothiazol-6-yl)-2-[(pyridin-4-ylmethyl)amino]-benzamide

10 N-(3,3-Dimethyl-1,1-dioxo-2,3-dihydro-1H-l\'benzo[d]isothiazol-6-yl)-2-[(pyridin-4-ylmethyl)-amino]benzamide (110 mg) was dissolved in DMF and added NaH (30
mg). The mix was stirred for 15 min then MeI (18 ul) was
added and stirred for 10 min. The Solvent was evaporated

15 and purified by preparative TLC (10% MeOH/EtOAc) to give the
product. M+H 438; Calc'd for C₂₂H₂₃N₅O₃S: 437.1.

The following compounds (Example 1013-1014) were synthesized by the method described above, unless specifically described.

1013) N-[3,3-Dimethyl-1,1-dioxo-2-(2-piperidin-1-yl-ethyl)2,3-dihydro-1H-1\(\lambda\)'-benzo[d]isothiazol-6-yl]-2[(pyridin-4-ylmethyl)-amino]-nicotinamide. M+H 535;
Calc'd for C28H4N6O3S: 534.

1014) N-[2-(2-Dimethylamino-ethyl)-3,3-dimethyl-1,1-dioxo2,3-dihydro-1H-l\(^1\)-benzo[d]isothiazol-6-yl]-2[(pyridin-4-ylmethyl)-amino]-nicotinamide. M+H 495;
Calc'd 494.

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## Example 1015

10 2-[(2,3-Dihydro-benzofuran-5-ylmethy1)-amino]-N-(1-Boc-4,4-dimethy1-1,2,3,4-tetrahydro-isoquinolin-7-y1)nicotinamide

M+H 529.4. Calc'd for 528.3.

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# Example 1016

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2-[(2,3-Dihydro-benzofuran-5-ylmethyl)-amino]-N-(4,4-dimethyl-1,2,3,4-tetrahydro-isoquinolin-7-yl)-nicotinamide

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M+H 429.2. Calc'd for 228.2.

# Example 1017

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2-[(2,3-Dihydro-benzofuran-5-ylmethyl)-amino]-N-[4pentafluoroethyl-3-(pyrrolidin-2-ylmethoxy)-ph nyl]nicotinamide

M+H 663.4. Calc'd for 662.3.

## Example 1018

N-(3,3-Dimethyl-2,3-dihydro-1H-indol-6-yl)-2-[(tetrahydropyran-4-ylmethyl)-amino]-nicotinamide

M+H 381.3, Calc'd for .

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## Example 1019

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N-(1-Acetyl-3,3-dimethyl-2,3-dihydro-1H-indol-6-yl)-2[(tetrahydro-pyran-4-ylmethyl)-amino]-nicotinamide

M+H 430. Calc'd for .

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# Example 1020

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# N-(4-Pentafluoroethyl-phenyl)-2-[(tetrahydro-pyran-4vlmethvl)-aminol-nicotinamide

M+H 432.2. Calc'd for .

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Although the pharmacological properties of the compounds of Formula I-XII vary with structural change, in general, activity possessed by compounds of Formula I-XII may be demonstrated in vivo. The pharmacological properties 10 of the compounds of this invention may be confirmed by a number of pharmacological in vitro assays. The exemplified pharmacological assays which follow have been carried out with the compounds according to the invention and their salts. Compounds of the present invention showed inhibition of KDR kinase at doses less than 50 uM.

### BIOLOGICAL EVALUATION

### **HUVEC Proliferation Assay**

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Human Umbilical Vein Endothelial cells are purchased from Clonetics, Inc., as cryopreserved cells harvested from a pool of donors. These cells, at passage 1, are thawed and expanded in EBM-2 complete medium, until passage 2 or 3. 25 The cells are trypsinized, washed in DMEM + 10% FBS + antibiotics, and spun at 1000 rpm for 10 min. Prior to centrifugation of the cells, a small amount is collected for a cell count. After centrifugation, the medium is discarded, and the cells are resuspended in the appropriate volume of DMEM + 10% FBS + antibiotics to achieve a 3.0 concentration of 3x105 cells/mL. Another cell count is performed to confirm the cell concentration. The cells are diluted to 3x104 cells/mL in DMEM + 10% FBS + antibiotics, and 100 µL of cells are added to a 96-well plate. The cells 35 are incubated at 37°C for 22 h.

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Prior to the completion of the incubation period, compound dilutions are prepared. Five-point, five-fold serial dilutions are prepared in DMSO, at concentrations 400-fold greater than the final concentrations desired. 2.5  $\mu$ L of each compound dilution are diluted further in a total of 1 mL DMEM + 10% FBS + antibiotics (400x dilution). Medium containing 0.25% DMSO is also prepared for the 0  $\mu$ M compound sample. At the 22-hour timepoint, the medium is removed from the cells, and 100  $\mu$ L of each compound dilution is added. The cells are incubated at 37°C for 2-3 h.

During the compound pre-incubation period, the growth factors are diluted to the appropriate concentrations. Solutions of DMEM  $\pm$  10% FBS  $\pm$  antibiotics, containing either VEGF or bFGF at the following concentrations: 50, 10, 2,

0.4, 0.08, and 0 ng/mL are prepared. For the compound-treated cells, solutions of VEGF at 550 ng/mL or bFGF at 220 ng/mL for 50 ng/mL or 20 ng/mL final concentrations, respectively, are prepared since 10 µL of each will be added to the cells (110 µL final volume). At the appropriate time after adding the compounds, the growth factors are added. VEGF is added to one set of plates, while bFGF is added to another set of plates. For the growth factor control curves, the media on wells B4-G6 of plates 1 and 2 are replaced with media containing VEGF or bFGF at the varying concentrations (50 - 0 ng/mL). The cells are incubated at 37°C for an additional 72 h.

At the completion of the 72 h incubation period, the medium is removed, and the cells are washed twice with PBS. After the second wash with PBS, the plates are tapped gently to remove excess PBS, and the cells are placed at -70°C for at least 30 min. The cells are thawed and analyzed using the CyQuant fluorescent dye (Molecular Probes C-7026), following the manufacturer's recommendations. The plates are read on a Victor/Wallac 1420 workstation at 485 nm/530

at a level below 50 nm.

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nm (excitation/emission). Raw data are collected and analyzed using a 4-parameter fit equation in XLFit.  $IC_{50}$  values are then determined.

Examples 4, 7, 20-21, 25-26, 28, 33, 67, 72(f-i, n-o), 5 78, 82, 84, 86, 94-95, 97-100, 105, 111-112, 115-118, 130, 133, 138, 140, 151, 154-156, 158-159, 165, 167, 169, 817, 826-829, 831-838, 840-844, 845, 847-851, 853, 855-860, 862, 864, 873, 900, 904-905, 916-917, 922-924, 942-944, 946, 951-952, 954-955, 963-964, 973, 977-978, 982, 985, 991, 995, 10 1000 and 1008 inhibited VEGF-stimulated HUVEC proliferation

### Angiogenesis Model

To determine the effects of the present compounds on angiogenesis in vivo, selective compounds are tested in the rat corneal neovascularization micropocket model or the angiogenesis assay of Passaniti, Lab. Invest., 67, 519-28 (1992).

### Rat Corneal Neovascularization Micropocket Model

In Life Aspects: Female Sprague Dawley rats weighing approximately 250 g were randomized into one of five treatment groups. Pretreatment with the vehicle or compound was administered orally, 24 h prior to surgery and continued once a day for seven additional days. On the day of surgery, the rats were temporarily anesthetized in an Isofluorane gas chamber (delivering 2.5 liters/min oxygen + 5% Isofluorane). An othoscope was then placed inside the mouth of the animal to visualize the vocal cords. A tipblunted wire was advanced in between the vocal cords and used as a guide for the placement of an endotracheal Teflon tube (Small Parts Inc. TFE-standard Wall R-SWTT-18). A volume-controlled ventilator (Harvard Apparatus, Inc. Model

overnight.

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683) was connected to the endotracheal tube to deliver a mixture of oxygen and 3% Isofluorane. Upon achieving deep anesthesia, the whiskers were cut short and the eye areas and eves gently washed with Betadine soap and rinsed with 5 sterile saline. The corneas were irrigated with one to two drops of Proparacaine HCl ophthalmic topical anesthetic solution (0.5%) (Bausch and Lomb Pharmaceuticals, Tampa FL). The rat was then positioned under the dissecting microscope and the corneal surface brought into focus. A vertical 10 incision was made on the midline of the cornea using a diamond blade knife. A pocket was created by using fine scissors to separate the connective tissue layers of the stroma, tunneling towards the limbus of the eve. The distance between the apex of the pocket and the limbus was 15 approximately 1.5 mm. After the pocket had been made, the soaked nitrocellulose disk filter (Gelman Sciences, Ann Arbor MI.) was inserted under the lip of the pocket. This surgical procedure was performed on both eyes. rHu-bFGF soaked disks were placed into the right eye, and the rHu-20 VEGF soaked disks were placed into the left eve. Vehicle soaked disks were placed in both eyes. The disk was pushed into position at the desired distance from the limbal vessels. Ophthalmic antibiotic ointment was applied to the eye to prevent drying and infection. After seven days, the 25 rats were euthanized by CO2 asphyxiation, and the eves enucleated. The retinal hemisphere of the eye was windowed to facilitate fixation, and the eye placed into formalin

Post Mortem Aspects: After twenty-four hours in

30 fixative, the corneal region of interest was dissected out
from the eye, using fine forceps and a razorblade. The
retinal hemisphere was trimmed off and the lens extracted
and discarded. The corneal dome was bisected and the
superfluous cornea trimmed off. The iris, conjunctiva and

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associated limbal glands were then carefully teased away. Final cuts were made to generate a square 3x3mm containing the disk, the limbus, and the entire zone of neovascularization.

5 Gross Image Recording: The corneal specimens were digitally photographed using a Sony CatsEye DKC5000 camera (A.G. Heinz, Irvine CA) mounted on a Nikon SMZ-U stereo microscope (A.G. Heinz). The corneas were submerged in distilled water and photographed via trans-illumination at approximately 5.0 diameters magnification.

Image analysis: Numerical endpoints were generated using digital micrographs collected from the whole mount corneas after trimming and were used for image analysis on the Metamorph image analysis system (Universal Imaging Corporation, West Chester PA). Three measurements were taken: Disk placement distance from the limbus, number of vessels intersecting a 2.0mm perpendicular line at the midpoint of the disk placement distance, and percent blood vessel area of the diffusion determined by thresholding.

### General Formulations:

0.1% BSA in PBS vehicle: 0.025 g of BSA was added to 25.0 ml of sterile lX phosphate buffered saline, gently shaken until fully dissolved, and filtered at 0.2  $\mu$ m. Individual 1.0 ml samples were aliquoted into 25 single use vials, and stored at -20 °C until use. For the rHu-bFGF disks, a vial of this 0.1% BSA solution was allowed to thaw at room temperature. Once thawed, 10  $\mu$ l of a 100 mM stock solution of DTT was added to the 1 ml BSA vial to yield a final concentration of 1 mM DTT in 0.1% BSA.

### 30 rHu-VEGF Dilutions:

Prior to the disk implant surgery, 23.8  $\mu l$  of the 0.1% BSA vehicle above was added to a 10  $\mu g$  rHu-VEGF lyophilized vial yielding a final concentration of 10  $\mu M$ .

rHu-bFGF: Stock concentration of 180 ng/μl:

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R&D rHu- bFGF: Added 139  $\mu$ l of the appropriate vehicle above to the 25  $\mu$ g vial lyophilized vial. 13.3  $\mu$ l of the [180 ng/ $\mu$ l] stock vial and added 26.6  $\mu$ l of vehicle to yield a final concentration of 3.75  $\mu$ M concentration.

5 Nitro-cellulose disk preparation: The tip of a 20-gauge needle was cut off square and beveled with emery paper to create a punch. This tip was then used to cut out ≅0.5mm diameter disks from a nitrocellulose filter paper sheet (Gelman Sciences). Prepared disks were then placed into 10 Eppendorf microfuge tubes containing solutions of either 0.1% BSA in PBS vehicle, 10 μM rHu-VEGF (R&D Systems, Minneapolis, MN), or 3.75 μM rHu-bFGF (R&D Systems, Minneapolis, MN) and allowed to soak for 45-60 min before use. Each nitrocellulose filter disk absorbs approximately 15 0.1 ul of solution.

In the rat micropocket assay, compounds of the present invention will inhibit angiogenesis at a dose of less than 50 mg/kg/day.

### Tumor model

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A431 cells (ATCC) are expanded in culture, harvested and injected subcutaneously into 5-8 week old female nude mice (CD1 nu/nu, Charles River Labs) (n=5-15). Subsequent administration of compound by oral gavage (10 - 200 mpk/dose) begins anywhere from day 0 to day 29 post tumor cell challenge and generally continues either once or twice a day for the duration of the experiment. Progression of tumor growth is followed by three dimensional caliper measurements and recorded as a function of time. Initial statistical analysis is done by repeated measures analysis of variance (RMANOVA), followed by Scheffe post hoc testing for multiple comparisons. Vehicle alone (Ora-Plus, pH 2.0) is the negative control. Compounds of the present invention are active at doses less than 150 mpk.

# ATTHECT TAKES

### Rat Adjuvant Arthritis Model:

The rat adjuvant arthritis model (Pearson, Proc. Soc. 5 Exp. Biol. 91, 95-101 (1956)) is used to test the antiarthritic activity of compounds of the formula I, or salts thereof. Adjuvant Arthritis can be treated using two different dosing schedules: either (i) starting time of immunization with adjuvant (prophylactic dosing); or from day 15 when the arthritic response is already established (therapeutic dosing). Preferably a therapeutic dosing schedule is used.

### Rat Carrageenan-induced Analgesia Test

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determined.

The rat carrageenan analgesia test was performed with materials, reagents and procedures essentially as described by Hargreaves, et al., (Pain, 32, 77 (1988)). Male Sprague-Dawley rats were treated as previously described for the Carrageenan Foot Pad Edema test. Three hours after the injection of the carrageenan, the rats were placed in a special plexiglass container with a transparent floor having a high intensity lamp as a radiant heat source, positionable under the floor. After an initial twenty minute period, thermal stimulation was begun on either the injected foot or on the contralateral uninjected foot. A photoelectric cell turned off the lamp and timer when light was interrupted by paw withdrawal. The time until the rat withdraws its foot was then measured. The withdrawal latency in seconds was determined for the control and drug-treated groups, and percent inhibition of the hyperalgesic foot withdrawal

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### Formulations

Also embraced within this invention is a class of pharmaceutical compositions comprising the active compounds of Formula I in association with one or more non-toxic. 5 pharmaceutically-acceptable carriers and/or diluents and/or adjuvants (collectively referred to herein as "carrier" materials) and, if desired, other active ingredients. The active compounds of the present invention may be administered by any suitable route, preferably in the form 10 of a pharmaceutical composition adapted to such a route, and in a dose effective for the treatment intended. The compounds and compositions of the present invention may, for example, be administered orally, mucosally, topically, rectally, pulmonarily such as by inhalation spray, or 15 parentally including intravascularly, intravenously, intraperitoneally, subcutaneously, intramuscularly intrasternally and infusion techniques, in dosage unit formulations containing conventional pharmaceutically acceptable carriers, adjuvants, and vehicles.

The pharmaceutically active compounds of this invention can be processed in accordance with conventional methods of pharmacy to produce medicinal agents for administration to patients, including humans and other mammals.

For oral administration, the pharmaceutical composition may be in the form of, for example, a tablet, capsule, suspension or liquid. The pharmaceutical composition is preferably made in the form of a dosage unit containing a particular amount of the active ingredient. Examples of such dosage units are tablets or capsules. For example, these may contain an amount of active ingredient

from about 1 to 2000 mg, preferably from about 1 to 500 mg or 5 to 1000 mg. A suitable daily dose for a human or other mammal may vary widely depending on the condition of

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the patient and other factors, but, once again, can be determined using routine methods.

The amount of compounds which are administered and the dosage regimen for treating a disease condition with the compounds and/or compositions of this invention depends on a variety of factors, including the age, weight, sex and medical condition of the subject, the type of disease, the severity of the disease, the route and frequency of administration, and the particular compound employed. Thus, the dosage regimen may vary widely, but can be determined routinely using standard methods. A daily dose of about 0.01 to 500 mg/kg, preferably between about 0.1 and about 50 mg/kg, and more preferably about 0.1 and about 20 mg/kg body weight may be appropriate. The daily dose can be administered in one to four doses per day.

For therapeutic purposes, the active compounds of this invention are ordinarily combined with one or more adjuvants appropriate to the indicated route of administration. If administered per os, the compounds may be admixed with lactose, sucrose, starch powder, cellulose esters of alkanoic acids, cellulose alkyl esters, talc, stearic acid, magnesium stearate, magnesium oxide, sodium and calcium salts of phosphoric and sulfuric acids, gelatin, acacia gum, sodium alginate, polyvinylpyrrolidone, and/or polyvinyl alcohol, and then tableted or encapsulated for convenient administration. Such capsules or tablets may contain a controlled-release formulation as may be provided in a dispersion of active compound in hydroxypropylmethyl cellulose.

In the case of psoriasis and other skin conditions, it may be preferable to apply a topical preparation of compounds of this invention to the affected area two to four times a day.

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from 0.1% to 1% of the formulation.

Formulations suitable for topical administration include liquid or semi-liquid preparations suitable for penetration through the skin (e.g., liniments, lotions, ointments, creams, or pastes) and drops suitable for administration to the eye, ear, or nose. A suitable topical dose of active ingredient of a compound of the invention is 0.1 mg to 150 mg administered one to four, preferably one or two times daily. For topical administration, the active ingredient may comprise from 0.001% to 10% w/w, e.g., from 1% to 2% by weight of the formulation, although it may comprise as much as 10% w/w, but preferably not more than 5% w/w, and more preferably

When formulated in an ointment, the active ingredients may be employed with either paraffinic or a water-miscible ointment base. Alternatively, the active ingredients may be formulated in a cream with an oil-in-water cream base. If desired, the aqueous phase of the cream base may include, for example at least 30% w/w of a polyhydric alcohol such as propylene glycol, butane-1,3-diol, mannitol, sorbitol, glycerol, polyethylene glycol and mixtures thereof. The topical formulation may desirably include a compound which enhances absorption or penetration of the active ingredient through the skin or other affected areas. Examples of such dermal penetration enhancers include dimethylsulfoxide and related analogs.

The compounds of this invention can also be administered by a transdermal device. Preferably transdermal administration will be accomplished using a patch either of the reservoir and porous membrane type or of a solid matrix variety. In either case, the active agent is delivered continuously from the reservoir or microcapsules through a membrane into the active agent permeable adhesive, which is in contact with the skin or mucosa of the recipient. If the

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active agent is absorbed through the skin, a controlled and predetermined flow of the active agent is administered to the recipient. In the case of microcapsules, the encapsulating agent may also function as the membrane.

The oily phase of the emulsions of this invention may be constituted from known ingredients in a known manner. While the phase may comprise merely an emulsifier, it may comprise a mixture of at least one emulsifier with a fat or an oil or with both a fat and an oil. Preferably, a hydrophilic emulsifier is included together with a

lipophilic emulsifier which acts as a stabilizer. It is also preferred to include both an oil and a fat. Together, the emulsifier(s) with or without stabilizer(s) make-up the so-called emulsifying wax, and the wax together with the oil and fat make up the so-called emulsifying ointment base which forms the oily dispersed phase of the cream formulations. Emulsifiers and emulsion stabilizers suitable for use in the formulation of the present invention include Tween 60, Span 80, cetostearyl alcohol, myristyl alcohol,

20 glyceryl monostearate, sodium lauryl sulfate, glyceryl distearate alone or with a wax, or other materials well known in the art.

The choice of suitable oils or fats for the formulation is based on achieving the desired cosmetic properties, since the solubility of the active compound in most oils likely to be used in pharmaceutical emulsion formulations is very low. Thus, the cream should preferably be a non-greasy, non-staining and washable product with suitable consistency to avoid leakage from tubes or other containers. Straight or branched chain, mono- or dibasic alkyl esters such as di-isoadipate, isocetyl stearate, propylene glycol diester of coconut fatty acids, isopropyl myristate, decyl oleate, isopropyl palmitate, butyl stearate, 2-ethylhexyl palmitate or a blend of branched

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chain esters may be used. These may be used alone or in combination depending on the properties required. Alternatively, high melting point lipids such as white soft paraffin and/or liquid paraffin or other mineral oils can be used.

Formulations suitable for topical administration to the eye also include eye drops wherein the active ingredients are dissolved or suspended in suitable carrier, especially an aqueous solvent for the active ingredients. The active ingredients are preferably present in such

The active ingredients are preferably present in such formulations in a concentration of 0.5 to 20%, advantageously 0.5 to 10% and particularly about 1.5% w/w.

Formulations for parenteral administration may be in the form of aqueous or non-aqueous isotonic sterile injection solutions or suspensions. These solutions and suspensions may be prepared from sterile powders or granules using one or more of the carriers or diluents mentioned for use in the formulations for oral administration or by using other suitable dispersing or wetting agents and suspending agents. The compounds may be dissolved in water, polyethylene glycol, propylene glycol, ethanol, corn oil, cottonseed oil, peanut oil, sesame oil, benzyl alcohol, sodium chloride, tragacanth gum, and/or various buffers. Other adjuvants and modes of administration are well and widely known in the pharmaceutical art. The active ingredient may also be administered by injection as a composition with suitable carriers including saline, dextrose, or water, or with cyclodextrin (ie. Captisol), cosolvent solubilization (ie. propylene glycol) or micellar solubilization (ie. Tween 80).

The sterile injectable preparation may also be a sterile injectable solution or suspension in a non-toxic parenterally acceptable diluent or solvent, for example as a solution in 1,3-butanediol. Among the acceptable vehicles

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and solvents that may be employed are water, Ringer's solution, and isotonic sodium chloride solution. In addition, sterile, fixed oils are conventionally employed as a solvent or suspending medium. For this purpose any bland fixed oil may be employed, including synthetic mono- or diglycerides. In addition, fatty acids such as oleic acid find use in the preparation of injectables.

For pulmonary administration, the pharmaceutical composition may be administered in the form of an aerosol or with an inhaler including dry powder aerosol.

Suppositories for rectal administration of the drug can be prepared by mixing the drug with a suitable non-irritating excipient such as cocoa butter and polyethylene glycols that are solid at ordinary temperatures but liquid at the rectal temperature and will therefore melt in the rectum and release the drug.

The pharmaceutical compositions may be subjected to conventional pharmaceutical operations such as sterilization and/or may contain conventional adjuvants, such as preservatives, stabilizers, wetting agents, emulsifiers, buffers etc. Tablets and pills can additionally be prepared with enteric coatings. Such compositions may also comprise adjuvants, such as wetting, sweetening, flavoring, and perfuming agents.

The foregoing is merely illustrative of the invention and is not intended to limit the invention to the disclosed compounds. Variations and changes which are obvious to one skilled in the art are intended to be within the scope and nature of the invention which are defined in the appended claims.

From the foregoing description, one skilled in the art can easily ascertain the essential characteristics of this invention, and without departing from the spirit and scope ANGUESA NAMES

thereof, can make various changes and modifications of the invention to adapt it to various usages and conditions.

All mentioned references, patents, applications and publications, are hereby incorporated by reference in their 5 entirety, as if here written.